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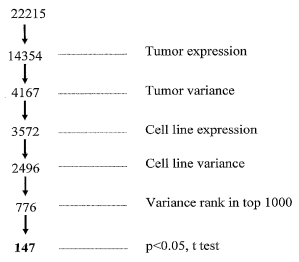
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[Continued on next page]

(54) Title: BIOMARKERS AND METHODS FOR DETERMINING SENSITIVITY TO EPIDERMAL GROWTH FACTOR RECEPTOR MODULATORS IN NON-SMALL CELL LUNG CANCER

Identification Scheme of Table 1 Biomarkers



(57) **Abstract:** EGFR biomarkers useful in a method for identifying a mammal that will respond therapeutically to a method of treating cancer comprising administering an EGFR modulator, wherein the method comprises (a) exposing a biological sample from the mammal to the EGFR modulator and (b) measuring in the biological sample the level of the at least one biomarker, wherein a difference in the level of the at least one biomarker measured in (b) compared to the level of the biomarker in a mammal that has not been exposed to the EGFR modulator indicates that the mammal will respond therapeutically to the method of treating cancer.





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BIOMARKERS AND METHODS FOR DETERMINING SENSITIVITY TO EPIDERMAL GROWTH FACTOR RECEPTOR MODULATORS IN NON-SMALL CELL LUNG CANCER

5 SEQUENCE LISTING:

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A compact disc labeled "Copy 1" contains the Sequence Listing as 10219 PCT.ST25.txt. The Sequence Listing is 1452 KB in size and was recorded March 24, 2005. The compact disk is 1 of 2 compact disks. A duplicate copy of the compact disc is labeled "Copy 2" and is 2 of 2 compact discs.

The compact disc and duplicate copy are identical and are hereby incorporated by reference into the present application.

FIELD OF THE INVENTION

The present invention relates generally to the field of pharmacogenomics, and more specifically to methods and procedures to determine drug sensitivity in patients to allow the identification of individualized genetic profiles which will aid in treating diseases and disorders.

BACKGROUND OF THE INVENTION:

Cancer is a disease with extensive histoclinical heterogeneity. Although conventional histological and clinical features have been correlated to prognosis, the same apparent prognostic type of tumors varies widely in its responsiveness to therapy and consequent survival of the patient.

New prognostic and predictive markers, which would facilitate an individualization of therapy for each patient, are needed to accurately predict patient response to treatments, such as small molecule or biological molecule drugs, in the clinic. The problem may be solved by the identification of new parameters that could better predict the patient's sensitivity to treatment. The classification of patient samples is a crucial aspect of cancer diagnosis and treatment. The association of a patient's response to a treatment with molecular and genetic markers can open up new opportunities for treatment development in non-responding patients, or distinguish a treatment's indication among other treatment choices because of higher confidence in the efficacy. Further, the pre-selection of patients who are likely to respond well to a medicine, drug, or combination therapy may reduce the number of patients needed in

a clinical study or accelerate the time needed to complete a clinical development program (M. Cockett et al., 2000, *Current Opinion in Biotechnology*, 11:602-609).

The ability to predict drug sensitivity in patients is particularly challenging because drug responses reflect not only properties intrinsic to the target cells, but also a host's metabolic properties. Efforts to use genetic imformation to predict drug sensitivity have primarily focused on individual genes that have broad effects, such as the multidrug resistance genes, *mdr1* and *mrp1* (P. Somneveld, 2000, *J. Intern. Med.*, 247:521-534).

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The development of microarray technologies for large scale characterization of gene mRNA expression pattern has made it possible to systematically search for molecular markers and to categorize cancers into distinct subgroups not evident by traditional histopathological methods (J. Khan et al., 1998, Cancer Res., 58:5009-5013; A.A. Alizadeh et al., 2000, Nature, 403:503-511; M. Bittner et al., 2000, Nature, 406:536-540; J. Khan et al., 2001, Nature Medicine, 7(6):673-679; and T.R. Golub et al., 1999, Science, 286:531-537; U. Alon et al., 1999, Proc. Natl. Acad. Sci. USA, 96:6745-6750). Such technologies and molecular tools have made it possible to monitor the expression level of a large number of transcripts within a cell population at any given time (see, e.g., Schena et al., 1995, Science, 270:467-470; Lockhart et al., 1996, Nature Biotechnology, 14:1675-1680; Blanchard et al., 1996, Nature Biotechnology, 14:1649; U.S. Patent No. 5,569,588 to Ashby et al.).

Recent studies demonstrate that gene expression information generated by microarray analysis of human tumors can predict clinical outcome (L.J. van't Veer et al., 2002, *Nature*, 415:530-536; T. Sorlie et al., 2001, *Proc. Natl. Acad. Sci. USA*, 98:10869-10874; M. Shipp et al., 2002, *Nature Medic ine*, 8(1):68-74: G.Glinsky et al., 2004, *The Journal of Clin. Invest.*, 113(6):913-923). These findings bring hope that cancer treatment will be vastly improved by better predicting the response of individual tumors to therapy.

Needed are new and alternative methods and procedures to determine drug sensitivity in patients to allow the development of individualized genetic profiles which are necessary to treat diseases and disorders based on patient response at a molecular level.

SUMMARY OF THE INVENTION:

The invention provides methods and procedures for determining patient sensitivity to one or more Epidermal Growth Factor Receptor (EGFR) modulators. The invention also provides methods of determining or predicting whether an individual requiring therapy for a disease state such as cancer will or will not respond to treatment, prior to administration of the treatment, wherein the treatment comprises of one or more EGFR modulators. The one or more EGFR modulators are compounds that can be selected from, for example, one or more EGFR-specific ligands, one or more small molecule EGFR inhibitors, or one or more EGFR binding monoclonal antibodies.

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In one aspect, the invention provides a method for identifying a mammal that will respond therapeutically to a method of treating cancer comprising administering of an EGFR modulator, wherein the method comprises: (a) measuring in the mammal the level of at least one biomarker selected from the biomarkers of Table 1; (b) exposing a biological sample from the mammal to the EGFR modulator; (c) following the exposing in step (b), measuring in said biological sample the level of the at least one biomarker, wherein a difference in the level of the at least one biomarker measured in step (c) compared to the level of the at least one biomarker measured in step (a) indicates that the mammal will respond therapeutically to the said method of treating cancer.

A difference in the level of the biomarker that is sufficient to indicate whether the mammal will or will not respond therapeutically to the method of treating cancer can be readily determined by one of skill in the art using known techniques. The increase or decrease in the level of the biomarker can be correlated to determine whether the difference is sufficient to identify a mammal that will respond therapeutically. The difference in the level of the biomarker that is sufficient can, in one aspect, be predetermined prior to determining whether the mammal will respond therapeutically to the treatment. In one aspect, the difference in the level of the biomarker is a difference in the mRNA level (measured, for example, by RT-PCT or a microarray), such as at least a two-fold difference, at least a three-fold difference, or at least a four-fold difference in the level of expression. In another aspect, the difference in the level of the biomarker is determined by IHC. In another aspect, the

difference in the level of the biomarker refers to a p-value of <0.05 in Anova analysis. In yet another aspect, the difference is determined in an ELISA assay.

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As used herein, respond therapeutically refers to the alleviation or abrogation of the cancer. This means that the life expectancy of an individual affected with the cancer will be increased or that one or more of the symptoms of the cancer will be reduced or ameliorated. The term encompasses a reduction in cancerous cell growth or tumor volume. Whether a mammal responds therapeutically can be measured by many methods well known in the art, such as PET imaging.

The mammal can be, for example, a human, rat, mouse, dog, rabbit, pig sheep, cow, horse, cat, primate, or monkey.

The method of the invention can be, for example, an *irz vitro* method wherein the step of measuring in the mammal the level of at least one biomarker comprises taking a biological sample from the mammal and then measuring the level of the at least one biomarker in the biological sample. The biological sample can comprise, for example, at least one of serum, whole fresh blood, peripheral blood mononuclear cells, frozen whole blood, fresh plasma, frozen plasma, urine, saliva, skin, hair follicle, bone marrow, or tumor tissue.

The level of the at least one biomarker can be, for example, the level of protein and/or mRNA transcript of the at least one biomarker.

In another aspect, the invention provides a method for identifying a mammal that will respond therapeutically to a method of treating cancer comprising administering an EGFR modulator, wherein the method comprises: (a) exposing a biological sample from the mammal to the EGFR modulator; (b) following the exposing of step (a), measuring in said biological sample the level of at least one biomarker selected from the biomarkers of Table 1, wherein a difference in the level of the at least one biomarker measured in step (b), compared to the level of the at least one biomarker in a mammal that has not been exposed to said EGFR modulator, indicates that the mammal will respond therapeutically to said method of treating cancer.

In yet another aspect, the invention provides a method for testing or predicting whether a mammal will respond therapeutically to a method of treating cancer comprising administering an EGFR modulator, wherein the method comprises: (a)

measuring in the mammal the level of at least one biomarker selected from the biomarkers of Table 1; (b) exposing the mammal to the EGFR modulator; (c) following the exposing of step (b), measuring in the mammal the level of the at least one biomarker, wherein a difference in the level of the at least one biomarker measured in step (c) compared to the level of the at least one biomarker measured in step (a) indicates that the mammal will respond therapeutically to said method of treating cancer.

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In another aspect, the invention provides a method for determining whether a compound inhibits EGFR activity in a mammal, comprising: (a) exposing the mammal to the compound; and (b) following the exposing of step (a), measuring in the mammal the level of at least one biomarker selected from the biomarkers of Table 1, wherein a difference in the level of said biomarker measured in step (b), compared to the level of the biomarker in a mammal that has not been exposed to said compound, indicates that the compound inhibits EGFR activity in the mammal.

In yet another aspect, the invention provides a method for determining whether a mammal has been exposed to a compound that inhibits EGFR activity, comprising (a) exposing the mammal to the compound; and (b) following the exposing of step (a), measuring in the mammal the level of at least one biomarker selected from the biomarkers of Table 1, wherein a difference in the level of said biomarker measured in step (b), compared to the level of the biomarker in a mammal that has not been exposed to said compound, indicates that the mammal has been exposed to a compound that inhibits EGFR activity.

In another aspect, the invention provides a method for determining whether a mammal is responding to a compound that inhibits EGFR activity, comprising (a) exposing the mammal to the compound; and (b) following the exposing of step (a), measuring in the mammal the level of at least one biomarker selected from the biomarkers of Table 1, wherein a difference in the level of the at least one biomarker measured in step (b), compared to the level of the at least one biomarker in a mammal that has not been exposed to said compound, indicates that the mammal is responding to the compound that inhibits EGFR activity.

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As used herein, "responding" encompasses responding by way of a biological and cellular response, as well as a clinical response (such as improved symptoms, a therapeutic effect, or an adverse event), in a mammal.

The invention also provides an isolated biomarker selected from the biomarkers of Table 1. The biomarkers of the invention comprise sequences selected from the nucleotide and amino acid sequences provided in Table 1 and the Sequence Listing, as well as fragments and variants thereof.

The invention also provides a biomarker set comprising two or more biomarkers selected from the biomarkers of Table 1.

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The invention also provides kits for determining or predicting whether a patient would be susceptible or resistant to a treatment that comprises one or more EGFR modulators. The patient may have a cancer or tumor such as, for example, a non-small cell lung cancer (NSCLC) or tumor.

In one aspect, the kit comprises a suitable container that comprises one or more specialized microarrays of the invention, one or more EGFR modulators for use in testing cells from patient tissue specimens or patient samples, and instructions for use. The kit may further comprise reagents or materials for monitoring the expression of a biomarker set at the level of mRNA or protein.

In another aspect, the invention provides a kit comprising two or more biomarkers selected from the biomarkers of Table 1.

In yet another aspect, the invention provides a kit comprising at least one of an antibody and a nucleic acid for detecting the presence of at least one of the biomarkers selected from the biomarkers of Table 1. In one aspect, the kit further comprises instructions for determining whether or not a mammal will respond therapeutically to a method of treating cancer comprising administering a compound that inhibits EGFR activity. In another aspect, the instructions comprise the steps of (a) measuring in the mammal the level of at least one biomarker selected from the biomarkers of Table 1, (b) exposing the mammal to the compound, (c) following the exposing of step (b), measuring in the mammal the level of the at least one biomarker measured in step (c) compared to the level of the at least one biomarker measured in step (a) indicates that the mammal will respond therapeutically to said method of treating cancer.

The invention also provides screening assays for determining if a patient will be susceptible or resistant to treatment with one or more EGFR modulators.

The invention also provides a method of monitoring the treatment of a patient having a disease, wherein said disease is treated by a method comprising administering one or more EGFR modulators.

The invention also provides individualized genetic profiles which are necessary to treat diseases and disorders based on patient response at a molecular level.

The invention also provides specialized microarrays, e.g., oligonucleotide microarrays or cDNA microarrays, comprising one or more biomarkers having expression profiles that correlate with either sensitivity or resistance to one or more EGFR modulators.

The invention also provides antibodies, including polyclonal or monoclonal, directed against one or more biomarkers of the invention.

The invention will be better understood upon a reading of the detailed description of the invention when considered in connection with the accompanying figures.

BRIEF DESCRIPTION OF THE FIGURES:

- FIG. 1 illustrates the scheme used for identifying the Table 1 biomarkers.
 - FIG. 2 illustrates the scheme used for identifying the Table 2 biomarkers.
 - FIG. 3 shows the mRNA levels of EGFR determined by expression profiling of fourteen NSCLC cell lines.
 - FIG. 4 illustrates the variance analysis of expression profiles.
 - FIG. 5 illustrates the variance metric distribution of probe sets for the adenocarcinoma tumors.
 - FIG. 6 illustrates the variance metric distribution of probe sets for the cell lines.
 - FIG. 7 illustrates the scoring of staining of a Calgranulin B IHC Assay.

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DETAILED DESCRIPTION OF THE INVENTION:

Identification of biomarkers that provide rapid and accessible readouts of efficacy, drug exposure, or clinical response is increasingly important in the clinical development of drug candidates. Embodiments of the invention include measuring changes in the levels of secreted proteins, or plasma biomarkers, which represent one category of biomarker. In one aspect, plasma samples, which represent a readily accessible source of material, serves a surrogate tissue for biomarker analysis.

The invention provides biomarkers that respond to the modulation of a specific signal transduction pathway and also correlate with EGFR modulator sensitivity or resistance. These biomarkers can be employed for predicting response to one or more EGFR modulators. In one aspect, the biomarkers of the invention are those provided in Table 1 and the Sequence Listing, including both polynucleotide and polypeptide sequences.

15 TABLE 1 - Biomarkers

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Unigene title and	Affymetrix Description	Affymetrix
SEQ ID NO:		Probe Set
S100A14: S100	gb:NM_020672.1 /DEF=Homo sapiens S100-	218677_at
calcium binding	type calcium binding protein A14 (LOC57402),	
protein A14	mRNA. /FEA=mRNA /GEN=LOC57402	
(LOC57402)	/PROD=S100-type calcium binding protein	
	A14 /DB_XREF=gi:10190711 /UG=Hs.288998	
SEQ ID NOS: 1	S100-type calcium binding protein A14	
(DNA) and 148	/FL=gb:NM_020672.1 gb:BC005019.1	
(amino acid)	gb:AY007220.1	
JTB: jumping	gb:AF151056.1 /DEF=Homo sapiens HSPC222	210434_x_at
translocation	mRNA, complete cds. /FEA=mRNA	
breakpoint	/PROD=HSPC222 /DB_XREF=gi:7106833	
(LOC10899)	/UG=Hs.323093 Homo sapiens, jumping	
	translocation breakpoint, clone MGC:10274,	
SEQ ID NOS: 2	mRNA, complete cds /FL=gb:AF151056.1	
(DNA) and 149		
(amino acid)		
CDH1: cadherin 1,	gb:NM_004360.1 /DEF=Homo sapiens	201131_s_at
type 1 preproprotein	cadherin 1, type 1, E-cadherin (epithelial)	
(LOC999)	(CDH1), mRNA. /FEA=mRNA /GEN=CDH1	
	/PROD=cadherin 1, type 1, E-cadherin	
SEQ ID NOS: 3	(epithelial) /DB_XREF=gi:4757959	
(DNA) and 150	/UG=Hs.194657 cadherin 1, type 1, E-cadherin	
(amino acid)	(epithelial) /FL=gb:L08599.1 gb:NM_004360.1	

CYR61: cysteine-	gb:NM_001554.1 /DEF=Homo sapiens	201289 at
rich, angiogenic	cysteine-rich, angiogenic inducer, 61 (CYR61),	
inducer, 61	mRNA. /FEA=mRNA /GEN=CYR61	
(LOC3491)	/PROD=cysteine-rich, angiogenic inducer, 61	
	/DB XREF=gi:4504612 /UG=Hs.8867	
SEQ ID NOS: 4	cysteine-rich, angiogenic inducer, 61	
(DNA) and 151	/FL=gb:BC001271.1 gb:U62015.1	
(amino acid)	gb:AF003594.1 gb:AF031385.1	
,	gb:NM 001554.1	
TGFBI: transforming	gb:NM_000358.1 /DEF=Homo sapiens	201506 at
growth factor, beta-	transforming growth factor, beta-induced, 68kD	-
induced, 68kDa	(TGFBI), mRNA. /FEA=mRNA	
(LOC7045)	/GEN=TGFBI /PROD=transforming growth	
	factor, beta-induced, 68kD	
SEQ ID NOS: 5	/DB_XREF=gi:4507466 /UG=Hs.118787	
(DNA) and 152	transforming growth factor, beta-induced, 68kD	
(amino acid)	/FL=gb:BC000097.1 gb:BC004972.1	
	gb:M77349.1 gb:NM_000358.1	
PSPHL:	Consensus includes gb:BF968134 /FEA=EST	212509 s at
phosphoserine	/DB_XREF=gi:12335349	Salaria Salaria
phosphatase-like	/DB_XREF=est:602269121F1	
(LOC8781)	/CLONE=IMAGE:4357349 /UG=Hs.250723	
	FK506 binding protein 12-rapamycin associated	
SEQ ID NOS: 6	protein 1	
(DNA) and 153		
(amino acid)		
DKK1: dickkopf	gb:NM_012242.1 /DEF=Homo sapiens	204602_at
homolog 1	dickkopf (Xenopus laevis) homolog 1 (DKK1),	
(LOC22943)	mRNA. /FEA=mRNA /GEN=DKK1	
	/PROD=dickkopf (Xenopus laevis) homolog 1	
SEQ ID NOS: 7	/DB_XREF=gi:7110718 /UG=Hs.40499	
(DNA) and 154	dickkopf (Xenopus laevis) homolog 1	
(amino acid)	/FL=gb:AF127563.1 gb:AF177394.1	
	gb:NM_012242.1	
FHL1: four and a half	gb:NM_001449.1 /DEF=Homo sapiens four	201540_at
LIM domains 1	and a half LIM domains 1 (FHL1), mRNA.	
(LOC2273)	/FEA=mRNA /GEN=FHL1 /PROD=four and a	
CEO ID MOG A	half LIM domains 1 /DB_XREF=gi:4503720	
SEQ ID NOS: 8	/UG=Hs.239069 four and a half LIM domains 1	
(DNA) and 155	/FL=gb:U29538.1 gb:U60115.1	
(amino acid)	gb:NM_001449.1	

SSR4: signal sequence receptor,	gb:NM_006280.1 /DEF=Homo sapiens signal sequence receptor, delta (translocon-associated	201004_at
delta (LOC6748)	protein delta) (SSR4), mRNA. /FEA=mRNA /GEN=SSR4 /PROD=signal sequence receptor,	
SEQ ID NOS: 9	delta/DB XREF=gi:5454089/UG=Hs.102135	
(DNA) and 156	signal sequence receptor, delta (translocon-	
(amino acid)	associated protein delta) /FL=gb:BC003371.1	
(3333335 33533)	gb:NM_006280.1	
S100A9: S100	gb:NM_002965.2 /DEF=Homo sapiens S100	203535_at
calcium-binding	calcium-binding protein A9 (calgranulin B)	_
protein A9	(S100A9), mRNA. /FEA=mRNA	
(LOC6280)	/GEN=S100A9 /PROD=S100 calcium-binding	
	protein A9 /DB_XREF=gi:9845520	
SEQ ID NOS: 10	/UG=Hs.112405 S100 calcium-binding protein	
(DNA) and 157	A9 (calgranulin B) /FL=gb:M26311.1	
(amino acid)	gb:NM_002965.2	
SFN: stratifin	Cluster Incl. X57348:H.sapiens mRNA (clone	33322_i_at
(LOC2810)	9112) /cds=(165,911) /gb=X57348 /gi=23939	
	/ug=Hs.184510 /len=1407	
SEQ ID NOS: 11		
(DNA) and 158		
(amino acid)		
F2RL1: coagulation	Consensus includes gb:BE965369 /FEA=EST	213506_at
factor II (thrombin)	/DB_XREF=gi:11769659	
receptor-like 1	/DB_XREF=est:601659282R1	
precursor (LOC2150)	/CLONE=IMAGE:3895653 /UG=Hs.168102	
GEO ID NIGG 13	Human proteinase activated receptor-2 mRNA,	,
SEQ ID NOS: 12	3UTR	
(DNA) and 159		
(amino acid) SPUVE: protease,	chilly 007172 1 /DEE—Homo conions	202459 -+
serine, 23 precursor	gb:NM_007173.1 /DEF=Homo sapiens protease, serine, 23 (SPUVE), mRNA.	202458_at
(LOC11098)	/FEA=mRNA /GEN=SPUVE /PROD=protease,	
(LOCITO)	serine, 23 /DB XREF=gi:6005881	
SEQ ID NOS: 13	/UG=Hs.325820 protease, serine, 23	
(DNA) and 160	/FL=gb:AL136914.1 gb:BC001278.1	
(amino acid)	gb:AF015287.1 gb:NM 007173.1	
	gb:AF193611.1	
AMIGO2:	Consensus includes gb:AC004010	222108 at
amphoterin induced	/DEF=Human BAC clone GS1-99H8	_
gene 2 (LOC347902)	/FEA=CDS /DB_XREF=gi:2781385	
	/UG=Hs.121520 Human BAC clone GS1-99H8	
SEQ ID NOS: 14		
(DNA) and 161		
(amino acid)		

KRT7: keratin 7	gb:BC002700.1 /DEF=Homo sapiens, Similar	209016 s at
(LOC3855)	to keratin 7, clone MGC:3625, mRNA,	209010_s_at
(200302)	complete cds. /FEA=mRNA /PROD=Similar to	
SEQ ID NOS: 15	keratin 7 /DB XREF=gi:12803726	
(DNA) and 162	/UG=Hs.23881 keratin 7 /FL=gb:BC002700.1	
(amino acid)	gb:NM 005556.1	
RPL13: ribosomal	Consensus includes gb:AW574664 /FEA=EST	212191 x at
protein L13	/DB_XREF=gi:7246203 /DB_XREF=est:UI-	212191_x_at
(LOC6137)	HF-BL0-abw-d-10-0-UI.s1	
	/CLONE=IMAGE:3057859 /UG=Hs.180842	
SEQ ID NOS: 16	ribosomal protein L13	
(DNA) and 163	Trooponial protein E15	
(amino acid)		
AF1Q: AF1Q protein	gb:BC006471.1 /DEF=Homo sapiens, ALL1-	211071 s at
(LOC10962)	fused gene from chromosome 1q, clone	2110/1_s_at
(20010302)	MGC:4013, mRNA, complete cds.	
SEQ ID NOS: 17	/FEA=mRNA /PROD=ALL1-fused gene from	
(DNA) and 164	chromosome 1q /DB XREF=gi:13623686	
(amino acid)	/FL=gb:BC006471.1	
COL6A2: alpha 2	gb:AY029208.1 /DEF=Homo sapiens type VI	209156 s at
type VI collagen	collagen alpha 2 chain precursor (COL6A2)	207130_s_at
isoform 2C2	mRNA, complete cds, alternatively spliced.	
precursor (LOC1292)	/FEA=mRNA /GEN=COL6A2 /PROD=type VI	
produisor (2001252)	collagen alpha 2 chain precursor	
SEQ ID NOS: 18	/DB XREF=gi:13603393 /UG=Hs.159263	
(DNA) and 165	collagen, type VI, alpha 2 /FL=gb:AY029208.1	
(amino acid)	go.111025200.1	
COL6A1: collagen,	Consensus includes gb:AA292373 /FEA=EST	213428 s at
type VI, alpha 1	/DB_XREF=gi:1940353	215 120_5_41
precursor (LOC1291)	/DB XREF=est:zt51a09.s1	
	/CLONE=IMAGE:725848 /UG=Hs.108885	
SEQ ID NOS: 19	collagen, type VI, alpha 1	
(DNA) and 166	5	
(amino acid)		
SLC38A2: solute	gb:NM 018573.1 /DEF=Homo sapiens	218041 x at
carrier family 38,	hypothetical protein PRO1068 (PRO1068),	
member 2	mRNA. /FEA=mRNA /GEN=PRO1068	
(LOC54407)	/PROD=hypothetical protein PRO1068	
, ,	/DB XREF=gi:8924006 /UG=Hs.321158	
SEQ ID NOS: 20	hypothetical protein PRO1068	
(DNA) and 167	/FL=gb:AF116620.1 gb:NM 018573.1	
(amino acid)		
		

PAPSS2: 3'-	gb:AF074331.1 /DEF=Homo sapiens PAPS	203060 s at
phosphoadenosine 5'-	synthetase-2 (PAPSS2) mRNA, complete cds.	203000_s_at
phosphosulfate	/FEA=mRNA/GEN=PAPSS2/PROD=PAPS	
synthase 2		
	synthetase-2 /DB_XREF=gi:5052074	
(LOC9060)	/UG=Hs.274230 3-phosphoadenosine 5-	
GEO TE MOS AL	phosphosulfate synthase 2 /FL=gb:AF150754.2	
SEQ ID NOS: 21	gb:AF313907.1 gb:AF091242.1	
(DNA) and 168	gb:NM_004670.1 gb:AF074331.1	
(amino acid)	gb:AF173365.1	
JAG1: jagged 1	gb:U73936.1 /DEF=Homo sapiens Jagged 1	209099_x_at
precursor (LOC182)	(HJ1) mRNA, complete cds. /FEA=mRNA	
	/GEN=HJ1 /PROD=Jagged 1	
SEQ ID NOS: 22	/DB XREF=gi:1695273 /UG=Hs.91143 jagged	
(DNA) and 169	1 (Alagille syndrome) /FL=gb:U61276.1	
(amino acid)	gb:U73936.1 gb:AF003837.1 gb:AF028593.1	
	gb:NM 000214.1	
RPS27L: ribosomal	gb:NM_015920.1 /DEF=Homo sapiens 40S	218007 s at
protein S27-like	ribosomal protein S27 isoform (LOC51065),	2,000,_5_4
protein (LOC51065)	mRNA. /FEA=mRNA /GEN=LOC51065	
protein (ECC31003)	/PROD=40S ribosomal protein S27 isoform	
SEQ ID NOS: 23	/DB XREF=gi:7705705 /UG=Hs.108957 40S	
(DNA) and 170	ribosomal protein S27 isoform	
(amino acid)	l	
(ammo acid)	/FL=gb:BC003667.1 gb:AF070668.1	
DAM montidal alvaina	gb:NM_015920.1	2022264
PAM: peptidylglycine	gb:NM_000919.1 /DEF=Homo sapiens	202336_s_at
alpha-amidating	peptidylglycine alpha-amidating	
monooxygenase	monooxygenase (PAM), mRNA. /FEA=mRNA	
isoform a,	/GEN=PAM /PROD=peptidylglycine alpha-	
preproprotein	amidating monooxygenase	
(LOC5066)	/DB_XREF=gi:4505602 /UG=Hs.83920	
	peptidylglycine alpha-amidating	
SEQ ID NOS: 24	monooxygenase /FL=gb:M37721.1	
(DNA) and 171	gb:NM_000919.1	
(amino acid)		
STAT1: signal	gb:NM_007315.1 /DEF=Homo sapiens signal	200887_s_at
transducer and	transducer and activator of transcription 1,	
activator of	91kD (STAT1), mRNA. /FEA=mRNA	
transcription 1	/GEN=STAT1 /PROD=signal transducer and	
isoform alpha	activator of transcription1, 91kD	
(LOC6772)	/DB_XREF=gi:6274551 /UG=Hs.21486 signal	
	transducer and activator of transcription 1,	
SEQ ID NOS: 25	91kD /FL=gb:M97935.1 gb:NM 007315.1	
(DNA) and 172	<u> </u>	
(amino acid)		
(DNA) and 172	71.17 80.1417/222.1 80.14141 00/212.1	

CTSB: cathepsin B	gb:NM_001908.1 /DEF=Homo sapiens	200839 s at
preproprotein	cathepsin B (CTSB), mRNA. /FEA=mRNA	200035_5_41
(LOC1508)	/GEN=CTSB /PROD=cathepsin B	
	/DB XREF=gi:4503138 /UG=Hs.297939	
SEQ ID NOS: 26	cathepsin B /FL=gb:M14221.1 gb:L16510.1	
(DNA) and 173	gb:NM 001908.1	
(amino acid)	g5.14141_001900.1	
POLR2L: DNA	gb:BC005903.1 /DEF=Homo sapiens,	211730 s at
directed RNA	polymerase (RNA) II (DNA directed)	211730_s_at
polymerase II	polypeptide L (7.6kD), clone MGC:14494,	
polymerase II polypeptide L	mRNA, complete cds. /FEA=mRNA	
(LOC5441)	/PROD=polymerase (RNA) II (DNA directed)	
(LOC5441)	polypeptide L(7.6kD) /DB_XREF=gi:13543491	
SEQ ID NOS: 27	polypeptide L(1.0kD)/DB_XKEF=gl.13343491 /FL=gb:BC005903.1	
(DNA) and 174	//L-go.bcoo5905.1	
(amino acid)		
ETV1: ets variant	Congangua includes ch.DE991500 /EDA- DCT	221011
	Consensus includes gb:BE881590 /FEA=EST /DB XREF=gi:10330366	221911_at
gene 1 (LOC2115)		
SEO ID NOS. 28	/DB_XREF=est:601490008F1	
SEQ ID NOS: 28	/CLONE=IMAGE:3892465 /UG=Hs.10684	
(DNA) and 175	Homo sapiens clone 24421 mRNA sequence	
(amino acid)	1.277.6.000004.1.77777777	201705
KRT18: keratin 18	gb:NM_000224.1 /DEF=Homo sapiens keratin	201596_x_at
(LOC3875)	18 (KRT18), mRNA. /FEA=mRNA	
GEO ID MOG GO	/GEN=KRT18 /PROD=keratin 18	
SEQ ID NOS: 29	/DB_XREF=gi:4557887 /UG=Hs.65114 keratin	
(DNA) and 176	18 /FL=gb:BC000698.1 gb:BC000180.2	
(amino acid)	gb:BC004253.1 gb:M26326.1	
	gb:NM_000224.1	
RPL29: ribosomal	Consensus includes gb:BF683426 /FEA=EST	213969_x_at
protein L29	/DB_XREF=gi:11968834	
(LOC6159)	/DB_XREF=est:602139603F1	
GTG TD 3.7GG 4.0	/CLONE=IMAGE:4300777 /UG=Hs.183698	
SEQ ID NOS: 30	ribosomal protein L29	
(DNA) and 177		
(amino acid)		
PYGB: brain	gb:NM_002862.1 /DEF=Homo sapiens	201481_s_at
glycogen	phosphorylase, glycogen; brain (PYGB),	
phosphorylase	nuclear gene encoding mitochondrial protein,	
(LOC5834)	mRNA. /FEA=mRNA /GEN=PYGB	
	/PROD=phosphorylase, glycogen; brain	
SEQ ID NOS: 31	/DB_XREF=gi:4506350 /UG=Hs.75658	
(DNA) and 178	phosphorylase, glycogen; brain	
(amino acid)	/FL=gb:U47025.1 gb:NM_002862.1	

ALCAM: activated	Consensus includes gb:AA156721 /FEA=EST	201952 at
leukocyte cell	/DB_XREF=gi:1728335	201932_at
adhesion molecule	/DB_XREF=est:zl18b04.s1	
(LOC214)	/CLONE=IMAGE:502255 /UG=Hs.10247	
(LOC214)		
SEQ ID NOS: 32	activated leucocyte cell adhesion molecule	
1 ~	/FL=gb:NM_001627.1 gb:L38608.1	
(DNA) and 179		
(amino acid)		
CTGF: connective	gb:M92934.1 /DEF=Human connective tissue	209101_at
tissue growth factor	growth factor, complete cds. /FEA=mRNA	
(LOC1490)	/PROD=connective tissue growth factor	
	/DB_XREF=gi:180923 /UG=Hs.75511	
SEQ ID NOS: 33	connective tissue growth factor	
(DNA) and 180	/FL=gb:M92934.1 gb:NM 001901.1	
(amino acid)		
UCHL1: ubiquitin	gb:NM_004181.1 /DEF=Homo sapiens	201387_s_at
carboxyl-terminal	ubiquitin carboxyl-terminal esterase L1	
esterase L1 (ubiquitin	(ubiquitin thiolesterase) (UCHL1), mRNA.	
thiolesterase)	/FEA=mRNA/GEN=UCHL1	
(LOC7345)	/PROD=ubiquitin carboxyl-terminal esterase	
	L1(ubiquitin thiolesterase)	
SEQ ID NOS: 34	/DB_XREF=gi:4759283 /UG=Hs.76118	
(DNA) and 181	ubiquitin carboxyl-terminal esterase L1	
(amino acid)	(ubiquitin thiolesterase) /FL=gb:BC000332.1	
(gb:BC005117.1 gb:NM 004181.1	
C14orf78:	Consensus includes gb:AI935123 /FEA=EST	212992 at
chromosome 14 open	/DB_XREF=gi:5673993	212992_at
reading frame 78	/DB_XREF=est:wp13h09.x1	
(LOC113 146)	/CLONE=IMAGE:2464769 /UG=Hs.57548	
(EGC113140)	ESTs	
SEQ ID NOS: 35	12019	
(DNA) and 182		
(amino acid)		
	Consequence in the state of DESCRIPTION DESCRIPTION	017700
PBEF: pre-B-cell	Consensus includes gb:BF575514 /FEA=EST	217738_at
colony-enhancing	/DB_XREF=gi:11649318	
factor isoform a	/DB_XREF=est:602133090F1	
(LOC10135)	/CLONE=IMAGE:4288079 /UG=Hs.239138	
	pre-B-cell colony-enhancing factor	
SEQ ID NOS: 36	/FL=gb:U02020.1 gb:NM_005746.1	
(DNA) and 183		
(amino acid)		

GNG11: guanine nucleotide binding protein (G protein), gamma 11 (LOC2791) SEQ ID NOS: 37 (DNA) and 184 (amino acid)	gb:NM_004126.1 /DEF=Homo sapiens guanine nucleotide binding protein 11 (GNG11), mRNA. /FEA=mRNA /GEN=GNG11 /PROD=guanine nucleotide binding protein 11 /DB_XREF=gi:4758447 /UG=Hs.83381 guanine nucleotide binding protein 11 /FL=gb:NM_004126.1 gb:U31384.1	204115_at
SERPINE2: plasminogen activator inhibitor type 1, member 2 (LOC5270) SEQ ID NOS: 38 (DNA) and 185 (amino acid)	Consensus includes gb:AL541302 /FEA=EST /DB_XREF=gi:12872241 /DB_XREF=est:AL541302 /CLONE=CS0DE006YI10 (5 prime) /UG=Hs.21858 trinucleotide repeat containing 3	212190_at
PTTG1IP: pituitary tumor-transforming gene 1 protein-interacting protein precursor (LOC754) SEQ ID NOS: 39 (DNA) and 186 (amino acid)	gb:NM_004339.2 /DEF=Homo sapiens pituitary tumor-transforming 1 interacting protein (PTTG1IP), mRNA. /FEA=mRNA /GEN=PTTG1IP /PROD=pituitary tumor-transforming protein1-interacting protein precursor /DB_XREF=gi:11038670 /UG=Hs.111126 pituitary tumor-transforming 1 interacting protein /FL=gb:NM_004339.2 gb:BC000415.1 gb:AF149785.1	200677_at
KRT19: keratin 19 (LOC3880) SEQ ID NOS: 40 (DNA) and 187 (amino acid)	gb:NM_002276.1 /DEF=Homo sapiens keratin 19 (KRT19), mRNA. /FEA=mRNA /GEN=KRT19 /PROD=keratin 19 /DB_XREF=gi:4504916 /UG=Hs.182265 keratin 19 /FL=gb:BC002539.1 gb:NM_002276.1	201650_at
SFN: stratifin (LOC2810) SEQ ID NOS: 41 (DNA) and 188 (amino acid)	Cluster Incl. X57348:H.sapiens mRNA (clone 9112) /cds=(165,911) /gb=X57348 /gi=23939 /ug=Hs.184510 /len=1407	33323_r_at
ICAM1: intercellular adhesion molecule 1 (LOC3383) SEQ ID NOS: 42 (DNA) and 189 (amino acid)	Consensus includes gb:AI608725 /FEA=EST /DB_XREF=gi:4617892 /DB_XREF=est:tw90b01.x1 /CLONE=IMAGE:2266921 /UG=Hs.168383 intercellular adhesion molecule 1 (CD54), human rhinovirus receptor /FL=gb:M24283.1 gb:J03132.1 gb:NM_000201.1	202637_s_at

SLCASA: solute carrier family 6 (neurotransmitter transporter, creatine), member 8 (SLCASA), mRNA. //EA=mRNA /GEN=SLC6A8 //PROD=solute carrier family 6 (neurotransmitter transporter, creatine), member 8 (JCC6535) SEQ ID NOS: 43 (DNA) and 190 (amino acid) ILS: interleukin 8 (LOC3576) SEQ ID NOS: 44 (DNA) and 191 (amino acid) (amino acid) (CSPG2: chondroitin sulfate proteoglycan 2 (versican) (LOC1462) (CSPG2: chondroitin sulfate proteoglycan 2 (versican) (CSPG2: mRNA. /GEN=mRNA /GEN=HRNA /GEN=BRNA /GEN=CSPG2 //PROD=chondroitin sulfate proteoglycan 2 (versican) (CSPG2: cathepsin C isoform a preproprotein (LOC1075) SEQ ID NOS: 46 (DNA) and 193 (amino acid) JTB: jumping translocation breakpoint (LOC10899) KRTS: kcratin 8 (LOC3856) KRTS: kcratin 8 (LOC3856) (DNA) and 194 (amino acid) KRTS: kcratin 8 (LOC3856) (DNOS: 48 (DNA) and 195 (gmn) acid) SEQ ID NOS: 48 (DNA) and 195 (gmn) acid) SEQ ID NOS: 48 (DNA) and 195 (gmn) acid) SEQ ID NOS: 48 (DNA) and 195 (gmn) acid) SEQ ID NOS: 49 (DNOS:	SLC6A8: solute	-1-NIM 005 (20.1 /DEE 11	200010
(neurotransmitter transporter, creatine), member 8 (SLC6A8), mRNA. /FEA=mRNA /GEN=SLC6A8 /PROD=solute carrier family 6 (neurotransmitter transporter, creatine), member 8 /DB_XREF=gi:5032096 /UG=Hs.187958 solute carrier family 6 (neurotransmitter transporter, creatine), member 8 /DB_XREF=gi:5032096 /UG=Hs.187958 solute carrier family 6 (neurotransmitter transporter, creatine), member 8 /DB_XREF=gi:5032096 /UG=Hs.187958 solute carrier family 6 (neurotransmitter transporter, creatine), member 8 /DB_XREF=gi:5032096 /UG=Hs.187958 solute carrier family 6 (neurotransmitter transporter, creatine), member 8 /DB_XREF=gi:5032096 /UG=Hs.187958 solute carrier family 6 (neurotransmitter transporter, creatine), member 8 /DB_XREF=gi:5032096 /UG=Hs.287958 solute carrier family 6 (neurotransmitter transporter, creatine), member 8 /DB_XREF=gi:5032096 /UG=Hs.032096 /UG=Hs.2879096 /UG=Hs.2879096 /UG=Hs.2879096 /UG=Hs.2879096 /UG=Hs.2879096 /UG=Hs.2879096 /UG=Hs.2879096 /UG=Hs.2879096 /UG=Hs.2879096 /UG=Hs.2879996 /UG=Hs.2879999 /UG=Hs.2879999 /UG=Hs.287999999999999999999999999999999999999			202219_at
transporter, creatine), member 8 (LOC6535) (LOC6535) (LOC6535) (ED NOS: 43 (DNA) and 190 (amino acid) II.8: interleukin 8 (LOC3576) (LOC3576) SEQ ID NOS: 44 (DNA) and 191 (amino acid) (SEQ ID NOS: 44 (DNA) and 191 (AD (SEP) (SE	· · · · · · · · · · · · · · · · · · ·		
member 8 (LOC6535) SEQ ID NOS: 43 (DNA) and 190 (amino acid) IL8: interleukin 8 (LOC3576) SEQ ID NOS: 44 (DNA) and 191 (amino acid) CSPG2: chondroitin sulfate proteoglycan 2 (versican) (CSPG2), mRNA. /FEA=mRNA /GEN=LSPSNA (SP) (CSPG2), mRNA. /FEA=mRNA (FEN=LSPS) (CSPG2), mRNA. /FEA=mRNA (FEN=CSPG2), mRNA. /FEA=mRNA (FEN=LSPS) (CSPG2), mRNA. /FEA=mRNA (FEN=CSPG2), mRNA. /FEA=mRNA (FEN=LSPS) (CSPG2), mRNA, complete cds. /FEA=mRNA (FEN=LSPS) (CSPG2), mRNA, complete cds. /FEA=mRNA (FEN=LSPS) (CSPG2), mRNA, complete cds. /FEA=mRNA (FENLLS) (CSPG2), mRNA			
CLOC6535 Creatine member 8 /DB_XREF=gi:5032096 / /UG=Hs.187958 solute carrier family 6 (neurotransmitter transporter, creatine) member 8 /FL=gb:131409.1 gb:NM_005629.1			
SEQ ID NOS: 43 (neurotransmitter transporter, creatine), member (20			
SEQ ID NOS: 43 (DNA) and 190 (amino acid) IL8: interleukin 8 (LOC3576) gb:AF043337.1 /DEF=Homo sapiens interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /FEA=mRNA /GEN=IL8 /PROD=interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /FEA=mRNA /GEN=IL8 /PROD=interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /FEA=mRNA /GEN=IL8 /PROD=interleukin 8 C-terminal variant (DS) /DB_XREF=gi:12641914 /UG=Hs.624 interleukin 8 /FL=gb:AF043337.1	(LOC6535)		
(DNA) and 190		/UG=Hs.187958 solute carrier family 6	
(amino acid) IL.8: interleukin 8 (LOC3576) gb:AF043337.1 //DEF=Homo sapiens interleukin 8 C-terminal variant (IL8) mRNA, complete cds. //FEA=mRNA //GEN=IL8 SEQ ID NOS: 44 (DNA) and 191 (amino acid) CSPG2: chondroitin sulfate proteoglycan 2 (versican) (LOC1462) (CSPG2), mRNA. //FEA=mRNA //GEN=IL8 SEQ ID NOS: 45 (DNA) and 192 (amino acid) CTSC: cathepsin C isoform a preproprotein (LOC1075) SEQ ID NOS: 46 (DNA) and 193 (amino acid) TB: jumping translocation breakpoint (LOC10899) SEQ ID NOS: 47 (DNA) and 194 (ACO10899) SEQ ID NOS: 47 (DNA) and 194 (COC10899) SEQ ID NOS: 47 (DNA) and 194 (COC10856) KRT8: keratin 8 (LOC3856) Richards C-terminal variant (IL8) mRNA, complete cds. //FEA=mRNA //GEN=IL8 (JOEA) Additional variant (JOEA) Additional variant (JOEHS-BLOO4239.1 //DEF=Homo sapiens cathepsin C (CTSC), mRNA. //FEA=mRNA (JOEN=CTSC //PROD=cathepsin C (JOEC1075) JOB_XREF=gi:45503140 //UG=Hs.10029 cathepsin C //FL=gb:NM_001814.1 201487_at 201487_a	SEQ ID NOS: 43	(neurotransmitter transporter, creatine), member	
II.8: interleukin 8	(DNA) and 190	8 /FL=gb:L31409.1 gb:NM 005629.1	
interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /FEA=mRNA /GEN=IL8 /PROD=interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /FEA=mRNA /GEN=IL8 /PROD=interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /FEA=mRNA /GEN=IL8 /PROD=interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /FEA=mRNA /GEN=IL8 /PROD=interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /FEA=mRNA /GEN=IL8 /PROD=interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /FEA=mRNA /GEN=IL8 /PROD=interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /FEA=mRNA /GEN=IL8 /PROD=interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /FEA=mRNA /GEN=IL8 /PROD=interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /FEA=mRNA /GEN=IL8 /PROD=interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /FEA=mRNA /GEN=IL8 /PROD=interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /FEA=mRNA /GEN=IL8 /PROD=interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /FEA=mRNA /GEN=IL8 /PROD=interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /FEA=mRNA /GEN=IL8 /PROD=interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /FEA=mRNA /GEN=IL8 /PROD=interleukin 8 C-terminal variant (IL8 /PROD=interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /FEA=mRNA /GEN=IL8 /PROD=interleukin 8 C-terminal variant (IL8	(amino acid)		
interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /FEA=mRNA /GEN=IL8 (DNA) and 191 (amino acid)	IL8: interleukin 8	gb:AF043337.1 /DEF=Homo sapiens	211506 s at
complete cds. /FEA=mRNA /GEN=IL8 /PROD=interleukin 8 C-terminal variant /DB_XREF=gi:12641914 /UG=Hs.624 interleukin 8 /FL=gb:AF043337.1 CSPG2: chondroitin sulfate proteoglycan 2 (versican) (LOC1462) (CSPG2), mRNA. /FEA=mRNA /GEN=CSPG2 /PROD=chondroitin sulfate proteoglycan 2 (versican) (CSPG2), mRNA. /FEA=mRNA /GEN=CSPG2 /PROD=chondroitin sulfate proteoglycan 2 (versican) (CSPG2), mRNA. /FEA=mRNA /GEN=CSPG2 /PROD=chondroitin sulfate proteoglycan 2 (versican) /FL=gb:NM_004385.1 CTSC: cathepsin C isoform a preproprotein (LOC1075) SEQ ID NOS: 46 (DNA) and 193 (amino acid) JTB: jumping translocation breakpoint (LOC10899) SEQ ID NOS: 47 (DNA) and 194 (amino acid) KRT8: keratin 8 (LOC3856) KRT8: keratin 8 (LOC3856) SEQ ID NOS: 48 (DNA) and 195 SEQ ID NOS: 48 (DNA) and 195 SEQ ID NOS: 48 (DNA) and 195	(LOC3576)		
SEQ ID NOS: 44 (DNA) and 191 (amino acid)			
(DNA) and 191	SEO ID NOS: 44		
Camino acid interleukin 8 /FL=gb:AF043337.1 gb:NM_004385.1 /DEF=Homo sapiens chondroitin sulfate proteoglycan 2 (versican) (LOC1462) (CSPG2), mRNA. /FEA=mRNA /GEN=CSPG2 /PROD=chondroitin sulfate proteoglycan 2 (versican) (CSPG2), mRNA. /FEA=mRNA /GEN=CSPG2 /PROD=chondroitin sulfate proteoglycan 2 (versican) /DB_XREF=gi:4758081 /UG=Hs.81800 chondroitin sulfate proteoglycan 2 (versican) /FL=gb:NM_004385.1 (CTSC: cathepsin C isoform a preproprotein (LOC1075) (DB_XREF=gi:4503140 /UG=Hs.10029 cathepsin C /FL=gb:NM_001814.1 /DEF=Homo sapiens cathepsin C /FL=gb:NM_001814.1 (CTSC) /FEA=mRNA /GEN=CTSC /PROD=cathepsin C /FL=gb:NM_001814.1 (CTSC) /FEA=mRNA /GEN=CTSC /PROD=cathepsin C /FL=gb:NM_001814.1 (CTSC) /FEA=mRNA	1 -		
CSPG2: chondroitin sulfate proteoglycan 2 (versican) (LOC1462) (versican) (LOC1462) SEQ ID NOS: 45 (DNA) and 192 (amino acid) CTSC: cathepsin C isoform a preproprotein (LOC1075) SEQ ID NOS: 46 (DNA) and 193 (amino acid) JTB: jumping translocation breakpoint (LOC10899) SEQ ID NOS: 47 (DNA) and 194 (ACC10899) SEQ ID NOS: 47 (DNA) and 194 (ACC10856) SEQ ID NOS: 47 (DNA) and 194 (ACC10856) KRT8: keratin 8 (LOC3856) SEQ ID NOS: 48 (DNA) and 195 SEQ ID NOS: 48 (DNA) and 195 SEQ ID NOS: 48 (LOC3856) S			
sulfate proteoglycan 2 (versican) (LOC1462) SEQ ID NOS: 45 (DNA) and 192 (amino acid) CTSC: cathepsin C isoform a preproprotein (LOC1075) SEQ ID NOS: 46 (DNA) and 193 (amino acid) SEQ ID NOS: 46 (DNA) and 193 (amino acid) SEQ ID NOS: 46 (DNA) and 193 (amino acid) JTB: jumping translocation breakpoint (LOC10899) SEQ ID NOS: 47 (DNA) and 194 (amino acid) SEQ ID NOS: 47 (DNA) and 194 (Amino acid) SEQ ID NOS: 47 (DNA) and 194 (Amino acid) KRT8: keratin 8 (LOC3856) SEQ ID NOS: 48 (DNA) and 195 SEQ ID NOS: 48 (DNA) and 195 SEQ ID NOS: 48 (LOC3856) SEQ ID NOS: 48 (DNA) and 195 SEQ ID NOS: 48 (DNA) and 195 SEQ ID NOS: 48 (LOC3856) SEQ ID NOS: 48 (LOC3856) SEQ ID NOS: 48 (LOG4239.1 /DEF=Human cytokeratin 8 mRNA, complete cds /FL=gb:BC004239.1 /DEF=Human cytokeratin 8 mRNA, complete cds /FEA=mRNA /PROD=cytokeratin 8 /DB_XREF=gi:1673574 /PGD=cytokeratin 8 /DB_XREF=gi:1673574 /PGD=cytokeratin 8 /DB_XREF=gi:1673574 /PGD=cytokeratin 8 /FL=gb:BC000654.1 (DNA) and 195 SEQ ID NOS: 48 (UG=Hs.242463 keratin 8 /FL=gb:BC000654.1 (DNA) and 195			204620+
(versican) (LOC1462) (CSPG2), mRNA. /FEA=mRNA /GEN=CSPG2 /PROD=chondroitin sulfate proteoglycan 2 (versican) /DB_XREF=gi:4758081 /UG=Hs.81800 chondroitin sulfate proteoglycan 2 (versican) /FL=gb:NM_004385.1 CTSC: cathepsin C gb:NM_001814.1 /DEF=Homo sapiens cathepsin C (CTSC), mRNA. /FEA=mRNA /GEN=CTSC /PROD=cathepsin C (DR) XREF=gi:4503140 /UG=Hs.10029 cathepsin C /FL=gb:NM_001814.1 SEQ ID NOS: 46 (DNA) and 193 (amino acid) JTB: jumping translocation breakpoint (LOC10899) SEQ ID NOS: 47 (DNA) and 194 (amino acid) KRT8: keratin 8 (LOC3856) KRT8: keratin 8 (LOC3856) REQ ID NOS: 48 (DNA) and 195 SEQ ID NOS: 48 (DNA) and 195			204620_s_at
GEN=CSPG2 /PROD=chondroitin sulfate proteoglycan 2 (versican)	1 2		
SEQ ID NOS: 45 (DNA) and 192 (amino acid) CTSC: cathepsin C isoform a preproprotein (LOC1075) SEQ ID NOS: 46 (DNA) and 193 (amino acid) SEQ ID NOS: 46 (DNA) and 193 (amino acid) JTB: jumping translocation breakpoint (LOC10899) SEQ ID NOS: 47 (DNA) and 194 (amino acid) SEQ ID NOS: 47 (DNA) and 194 (amino acid) SEQ ID NOS: 47 (DNA) and 194 (amino acid) SEQ ID NOS: 47 (DNA) and 194 (amino acid) SEQ ID NOS: 47 (DNA) and 194 (amino acid) SEQ ID NOS: 47 (DNA) and 194 (amino acid) KRT8: keratin 8 (LOC3856) SEQ ID NOS: 48 (DNA) and 195	(versican) (LOC1462)		
(DNA) and 192 (amino acid) (DB_XREF=gi:4758081 /UG=Hs.81800 chondroitin sulfate proteoglycan 2 (versican) /FL=gb:NM_004385.1 CTSC: cathepsin C isoform a preproprotein (LOC1075) SEQ ID NOS: 46 (DNA) and 193 (amino acid) JTB: jumping translocation breakpoint (LOC10899) SEQ ID NOS: 47 (DNA) and 194 (amino acid) KRT8: keratin 8 (LOC3856) KRT8: keratin 8 (DNA) and 195 SEQ ID NOS: 48 (DNA) and 195 SEQ ID NOS: 48 (DNA) and 194 (amino acid) KRT8: keratin 8 (LOC3856) SDB_XREF=gi:4758081 /UG=Hs.81800 chondroitin sulfate proteoglycan 2 (versican) /FL=gb:NM_001814.1 /DEF=Homo sapiens CATHEPSINA /PROD=jumping translocation breakpoint /DB_XREF=gi:13278986 /UG=Hs.323093 FROD=jumping translocation breakpoint, clone MGC:10274, mRNA, complete cds /FL=gb:BC004239.1 KRT8: keratin 8 (BDA) RREF=gi:1673574 SEQ ID NOS: 48 (DNA) and 195	GDG D MGG 45		
chondroitin sulfate proteoglycan 2 (versican) //FL=gb:NM_004385.1 CTSC: cathepsin C isoform a preproprotein (LOC1075) SEQ ID NOS: 46 (DNA) and 193 (amino acid) JTB: jumping translocation breakpoint (LOC10899) SEQ ID NOS: 47 (DNA) and 194 (amino acid) KRT8: keratin 8 (LOC3856) KEQ ID NOS: 48 (DNA) and 195 SEQ ID NOS: 48 (LOC3856) Chondroitin sulfate proteoglycan 2 (versican) //FL=gb:NM_004385.1 gb:NM_001814.1 /DEF=Homo sapiens cathepsin C (CTSC), mRNA. /FEA=mRNA //GEN=CTSC //ROD=cathepsin C //DB_XREF=gi:4503140 //UG=Hs.10029 cathepsin C //FL=gb:NM_001814.1 SEQ ID NOS: 46 (DNA) and 193 (amino acid) gb:BC004239.1 //DEF=Homo sapiens, jumping translocation breakpoint, clone MGC:10274, mRNA, complete cds. //FEA=mRNA //PROD=jumping translocation breakpoint //DB_XREF=gi:13278986 //UG=Hs.323093 Homo sapiens, jumping translocation breakpoint, clone MGC:10274, mRNA, complete cds //FL=gb:BC004239.1 KRT8: keratin 8 (B:U76549.1 //DEF=Human cytokeratin 8 //PROD=cytokeratin 8 //DB_XREF=gi:1673574 //PROD=cytokeratin 8 //DB_XREF=gi:1673574 //UG=Hs.242463 keratin 8 //FL=gb:BC000654.1 (DNA) and 195 SEQ ID NOS: 48 (DNA) and 195 SEQ ID NOS: 48 (DNA) and 195 SEQ ID NOS: 48 (DNA) and 195	_	, ,	
CTSC: cathepsin C gb:NM_004385.1 gb:NM_001814.1 /DEF=Homo sapiens cathepsin C (CTSC), mRNA. /FEA=mRNA /GEN=CTSC /PROD=cathepsin C (DOC1075) /DB_XREF=gi:4503140 /UG=Hs.10029 cathepsin C /FL=gb:NM_001814.1 SEQ ID NOS: 46 (DNA) and 193 (amino acid) JTB: jumping translocation breakpoint (LOC10899) gb:BC004239.1 /DEF=Homo sapiens, jumping translocation breakpoint (LOC10899) mRNA, complete cds. /FEA=mRNA /PROD=jumping translocation breakpoint /DB_XREF=gi:13278986 /UG=Hs.323093 Homo sapiens, jumping translocation breakpoint (DNA) and 194 (amino acid) breakpoint, clone MGC:10274, mRNA, complete cds /FL=gb:BC004239.1 KRT8: keratin 8 (BCC3856) gb:U76549.1 /DEF=Human cytokeratin 8 209008_x_at mRNA, complete cds. /FEA=mRNA /PROD=cytokeratin 8 /DB_XREF=gi:1673574 SEQ ID NOS: 48 (UG=Hs.242463 keratin 8 /FL=gb:BC000654.1 (DNA) and 195 gb:U76549.1 gb:M34225.1 gb:M26324.1		, = •	
CTSC: cathepsin C isoform a preproprotein (LOC1075)	(amino acid)		
isoform a preproprotein (LOC1075) SEQ ID NOS: 46 (DNA) and 193 (amino acid) JTB: jumping translocation breakpoint (LOC10899) SEQ ID NOS: 47 (DNA) and 194 (amino acid) KRT8: keratin 8 (LOC3856) KRT8: keratin 8 (LOC3856) SEQ ID NOS: 48 (DNA) and 195 cathepsin C (CTSC), mRNA. /FEA=mRNA /GEN=CTSC /PROD=cathepsin C /DB_XREF=gi:4503140 /UG=Hs.10029 cathepsin C /FL=gb:NM_001814.1 SEQ ID NOS: 46 (DNA) and 193 (amino acid) gb:BC004239.1 /DEF=Homo sapiens, jumping translocation breakpoint, clone MGC:10274, mRNA, complete cds. /FEA=mRNA /PROD=jumping translocation breakpoint, clone MGC:10274, mRNA, complete cds /FL=gb:BC004239.1 KRT8: keratin 8 (LOC3856) KRT8: keratin 8 (LOC3856) SEQ ID NOS: 48 (LOC3856) SEQ ID NOS: 48 (LOC3856) Gathepsin C (CTSC), mRNA. /FEA=mRNA /PROD=cytokeratin 8 /DB_XREF=gi:1673574 /PROD=cytokeratin 8 /DB_XREF=gi:1673574 /UG=Hs.242463 keratin 8 /FL=gb:BC000654.1 (DNA) and 195 Gathepsin C (CTSC), mRNA. /FEA=mRNA /PROD=cytokeratin 8 /DB_XREF=gi:1673574 /UG=Hs.242463 keratin 8 /FL=gb:BC000654.1 gb:U76549.1 gb:M34225.1 gb:M26324.1		/FL=gb:NM_004385.1	
Description	CTSC: cathepsin C	gb:NM_001814.1 /DEF=Homo sapiens	201487_at
(LOC1075) /DB_XREF=gi:4503140 /UG=Hs.10029 cathepsin C /FL=gb:NM_001814.1 SEQ ID NOS: 46 (DNA) and 193 (amino acid) JTB: jumping translocation breakpoint (LOC10899) SEQ ID NOS: 47 (DNA) and 194 (amino acid) KRT8: keratin 8 (LOC3856) KRT8: keratin 8 (LOC3856) /DB_XREF=gi:4503140 /UG=Hs.10029 cathepsin C /FL=gb:NM_001814.1 gb:BC004239.1 /DEF=Homo sapiens, jumping translocation breakpoint, clone MGC:10274, mRNA /PROD=jumping translocation breakpoint /DB_XREF=gi:13278986 /UG=Hs.323093 Homo sapiens, jumping translocation breakpoint, clone MGC:10274, mRNA, (amino acid) kRT8: keratin 8 (LOC3856) gb:U76549.1 /DEF=Human cytokeratin 8 /PROD=cytokeratin 8 /DB_XREF=gi:1673574 /PROD=cytokeratin 8 /FL=gb:BC000654.1 (gb:U76549.1 gb:M34225.1 gb:M26324.1	isoform a	cathepsin C (CTSC), mRNA. /FEA=mRNA	
cathepsin C /FL=gb:NM_001814.1 SEQ ID NOS: 46 (DNA) and 193 (amino acid) JTB: jumping translocation breakpoint, clone MGC:10274, breakpoint (LOC10899) SEQ ID NOS: 47 (DNA) and 194 (amino acid) KRT8: keratin 8 (LOC3856) KRT8: keratin 8 (LOC3856) SEQ ID NOS: 48 (DNA) and 195 Cathepsin C /FL=gb:NM_001814.1 gb:BC004239.1 /DEF=Homo sapiens, jumping translocation breakpoint, clone MGC:10274, mRNA, complete cds. /FEA=mRNA (JEF) Cathepsin C /FL=gb:NM_001814.1 SEQ ID NOS: 46 (DNA) and 193 gb:BC004239.1 /DEF=Homo sapiens, jumping translocation breakpoint /PROD=jumping translocation breakpoint /DB_XREF=gi:13278986 /UG=Hs.323093 Homo sapiens, jumping translocation breakpoint, clone MGC:10274, mRNA, complete cds /FL=gb:BC004239.1 KRT8: keratin 8 (LOC3856) gb:U76549.1 /DEF=Human cytokeratin 8 /PROD=cytokeratin 8 /DB_XREF=gi:1673574 /PROD=cytokeratin 8 /DB_XREF=gi:1673574 /UG=Hs.242463 keratin 8 /FL=gb:BC000654.1 gb:U76549.1 gb:M34225.1 gb:M26324.1	preproprotein	/GEN=CTSC /PROD=cathepsin C	
cathepsin C /FL=gb:NM_001814.1 SEQ ID NOS: 46 (DNA) and 193 (amino acid) JTB: jumping translocation breakpoint (LOC10899) SEQ ID NOS: 47 (DNA) and 194 (amino acid) KRT8: keratin 8 (LOC3856) KRT8: keratin 8 (LOC3856) SEQ ID NOS: 48 (DNA) and 195 Cathepsin C /FL=gb:NM_001814.1 gb:BC004239.1 /DEF=Homo sapiens, jumping translocation breakpoint, clone MGC:10274, mRNA, complete cds. /FEA=mRNA /PROD=jumping translocation breakpoint /DB_XREF=gi:13278986 /UG=Hs.323093 Homo sapiens, jumping translocation breakpoint, clone MGC:10274, mRNA, complete cds /FL=gb:BC004239.1 KRT8: keratin 8 (LOC3856) gb:U76549.1 /DEF=Human cytokeratin 8 /PROD=cytokeratin 8 /DB_XREF=gi:1673574 /PROD=cytokeratin 8 /DB_XREF=gi:1673574 /UG=Hs.242463 keratin 8 /FL=gb:BC000654.1 gb:U76549.1 gb:M34225.1 gb:M26324.1	(LOC1075)	/DB XREF=gi:4503140 /UG=Hs.10029	
SEQ ID NOS: 46 (DNA) and 193 (amino acid) JTB: jumping translocation breakpoint (LOC10899) JRCD="index"		cathepsin C /FL=gb:NM 001814.1	
(DNA) and 193 (amino acid) JTB: jumping	SEQ ID NOS: 46	_	
(amino acid)gb:BC004239.1 /DEF=Homo sapiens, jumping translocation210927_x_atbreakpoint (LOC10899)mRNA, complete cds. /FEA=mRNA210927_x_atSEQ ID NOS: 47 (DNA) and 194 (amino acid)homo sapiens, jumping translocation breakpoint, clone MGC:10274, mRNA, complete cds /FL=gb:BC004239.1209008_x_atKRT8: keratin 8 (LOC3856)gb:U76549.1 /DEF=Human cytokeratin 8 /PROD=cytokeratin 8 /PROD=cytokeratin 8 /PROD=cytokeratin 8 /FL=gb:BC000654.1209008_x_at(DNA) and 195gb:U76549.1 gb:M34225.1 gb:M26324.1			
JTB: jumping translocation breakpoint, clone MGC:10274, mRNA, complete cds. /FEA=mRNA (LOC10899) /PROD=jumping translocation breakpoint /DB_XREF=gi:13278986 /UG=Hs.323093 Homo sapiens, jumping translocation breakpoint (DNA) and 194 breakpoint, clone MGC:10274, mRNA, complete cds /FL=gb:BC004239.1 KRT8: keratin 8 (LOC3856) /PROD=cytokeratin 8 /DB_XREF=gi:1673574 /PROD=cytokeratin 8 /DB_XREF=gi:1673574 /UG=Hs.242463 keratin 8 /FL=gb:BC000654.1 gb:U76549.1 gb:M34225.1 gb:M26324.1	()		
translocation breakpoint, clone MGC:10274, mRNA, complete cds. /FEA=mRNA (LOC10899) /PROD=jumping translocation breakpoint /DB_XREF=gi:13278986 /UG=Hs.323093 SEQ ID NOS: 47 Homo sapiens, jumping translocation breakpoint, clone MGC:10274, mRNA, complete cds /FL=gb:BC004239.1 KRT8: keratin 8 gb:U76549.1 /DEF=Human cytokeratin 8 gb:U76549.1 /DEF=Human cytokeratin 8 /PROD=cytokeratin 8 /PROD=cytokeratin 8 /PROD=cytokeratin 8 /PROD=cytokeratin 8 /PROD=cytokeratin 8 /FL=gb:BC000654.1 gb:U76549.1 gb:M34225.1 gb:M26324.1		gh:BC004239 1 /DEF=Homo saniens jumning	210927 x at
breakpoint (LOC10899) mRNA, complete cds. /FEA=mRNA /PROD=jumping translocation breakpoint /DB_XREF=gi:13278986 /UG=Hs.323093 Homo sapiens, jumping translocation breakpoint, clone MGC:10274, mRNA, (amino acid) krts: keratin 8 (LOC3856) gb:U76549.1 /DEF=Human cytokeratin 8 /PROD=cytokeratin 8 /DB_XREF=gi:1673574 /PROD=cytokeratin 8 /FL=gb:BC000654.1 gb:U76549.1 gb:M34225.1 gb:M26324.1			210721_K_at
(LOC10899) /PROD=jumping translocation breakpoint /DB_XREF=gi:13278986 /UG=Hs.323093 SEQ ID NOS: 47 Homo sapiens, jumping translocation (DNA) and 194 breakpoint, clone MGC:10274, mRNA, (amino acid) complete cds /FL=gb:BC004239.1 KRT8: keratin 8 gb:U76549.1 /DEF=Human cytokeratin 8 (LOC3856) mRNA, complete cds. /FEA=mRNA /PROD=cytokeratin 8 /DB_XREF=gi:1673574 /UG=Hs.242463 keratin 8 /FL=gb:BC000654.1 (DNA) and 195 gb:U76549.1 gb:M34225.1 gb:M26324.1			
DB_XREF=gi:13278986 /UG=Hs.323093 Homo sapiens, jumping translocation breakpoint, clone MGC:10274, mRNA, complete cds /FL=gb:BC004239.1 Example 10		_	
SEQ ID NOS: 47 Homo sapiens, jumping translocation (DNA) and 194 breakpoint, clone MGC:10274, mRNA, (amino acid) complete cds /FL=gb:BC004239.1 KRT8: keratin 8 gb:U76549.1 /DEF=Human cytokeratin 8 (LOC3856) mRNA, complete cds. /FEA=mRNA /PROD=cytokeratin 8 /DB_XREF=gi:1673574 SEQ ID NOS: 48 /UG=Hs.242463 keratin 8 /FL=gb:BC000654.1 (DNA) and 195 gb:U76549.1 gb:M34225.1 gb:M26324.1	(100100)		
(DNA) and 194 breakpoint, clone MGC:10274, mRNA, (amino acid) complete cds /FL=gb:BC004239.1 KRT8: keratin 8 gb:U76549.1 /DEF=Human cytokeratin 8 209008_x_at (LOC3856) mRNA, complete cds. /FEA=mRNA /PROD=cytokeratin 8 /DB_XREF=gi:1673574 SEQ ID NOS: 48 /UG=Hs.242463 keratin 8 /FL=gb:BC000654.1 (DNA) and 195 gb:U76549.1 gb:M34225.1 gb:M26324.1	SEO ID NOS. 47		
(amino acid) complete cds /FL=gb:BC004239.1 KRT8: keratin 8 gb:U76549.1 /DEF=Human cytokeratin 8 209008_x_at (LOC3856) mRNA, complete cds. /FEA=mRNA /PROD=cytokeratin 8 /DB_XREF=gi:1673574 SEQ ID NOS: 48 /UG=Hs.242463 keratin 8 /FL=gb:BC000654.1 (DNA) and 195 gb:U76549.1 gb:M34225.1 gb:M26324.1		1 ,0 1	
KRT8: keratin 8 gb:U76549.1 /DEF=Human cytokeratin 8 209008_x_at			
(LOC3856) mRNA, complete cds. /FEA=mRNA /PROD=cytokeratin 8 /DB_XREF=gi:1673574 /UG=Hs.242463 keratin 8 /FL=gb:BC000654.1 gb:U76549.1 gb:M34225.1 gb:M26324.1			200000
/PROD=cytokeratin 8 /DB_XREF=gi:1673574 SEQ ID NOS: 48 /UG=Hs.242463 keratin 8 /FL=gb:BC000654.1 (DNA) and 195 gb:U76549.1 gb:M34225.1 gb:M26324.1			209008_x_at
SEQ ID NOS: 48 /UG=Hs.242463 keratin 8 /FL=gb:BC000654.1 gb:U76549.1 gb:M34225.1 gb:M26324.1	(LOC3856)	· · · · · · · · · · · · · · · · · · ·	
(DNA) and 195 gb:U76549.1 gb:M34225.1 gb:M26324.1			
	_	i e	
(amino acid) gb:NM_002273.1	` ′		
	(amino acid)	gb:NM_002273.1	

UGDH: UDP-glucose	gb:NM_003359.1 /DEF=Homo sapiens UDP-	203343 at
dehydrogenase	guinni 003339.17DET—Hollio Sapielis ODF-	203343_at
	glucose dehydrogenase (UGDH), mRNA.	
(LOC7358)	/FEA=mRNA /GEN=UGDH /PROD=UDP-	
	glucose dehydrogenase	
SEQ ID NOS: 49	/DB_XREF=gi:4507812 /UG=Hs.28309 UDP-	
(DNA) and 196	glucose dehydrogenase /FL=gb:AF061016.1	
(amino acid)	gb:NM_003359.1	
TXNIP: thioredoxin	Consensus includes gb:AA812232 /FEA=EST	201008_s_at
interacting protein	/DB_XREF=gi:2881843	
(LOC10628)	/DB_XREF=est:ob84h09.s1	
	/CLONE=IMAGE:1338113 /UG=Hs.179526	
SEQ ID NOS: 50	upregulated by 1,25-dihydroxyvitamin D-3	
(DNA) and 197	/FL=gb:NM_006472.1 gb:S73591.1	
(amino acid)		
CTSB: cathepsin B	gb:NM_001908.1 /DEF=Homo sapiens	200838 at
preproprotein	cathepsin B (CTSB), mRNA. /FEA=mRNA	_
(LOC1508)	/GEN=CTSB /PROD=cathepsin B	
	/DB XREF=gi:4503138 /UG=Hs.297939	
SEQ ID NOS: 51	cathepsin B /FL=gb:M14221.1 gb:L16510.1	
(DNA) and 198	gb:NM 001908.1	
(amino acid)	2	
CSPG2: chondroitin	Consensus includes gb:BF218922 /FEA=EST	221731 x at
sulfate proteoglycan 2	/DB XREF=gi:11112418	
(versican) (LOC1462)	/DB XREF=est:601885091F1	
	/CLONE=IMAGE:4103447 /UG=Hs.81800	
SEQ ID NOS: 52	chondroitin sulfate proteoglycan 2 (versican)	
(DNA) and 199	1 2 3 3 4 4 4	
(amino acid)		
ANXA10: annexin	gb:AF196478.1 /DEF=Homo sapiens annexin	210143 at
A10 (LOC11199)	14 (ANX14) mRNA, complete cds.	
	/FEA=mRNA /GEN=ANX14 /PROD=annexin	
SEQ ID NOS: 53	14 /DB_XREF=gi:6274496 /UG=Hs.188401	
(DNA) and 200	annexin A10 /FL=gb:AF196478.1	
(amino acid)	gb:NM 007193.2	
SAT:	gb:M55580.1 /DEF=Human	210592 s at
spermidine/spermine	spermidinespermine N1-acetyltransferase	210072_5_at
N1-acetyltransferase	mRNA, complete cds. /FEA=mRNA	
(LOC6303)	/GEN=spermidinespermine N1-	
(1000000)	acetyltransferase /PROD=spermidinespermine	
SEQ ID NOS: 54	N1-acetyltransferase /DB XREF=gi:338335	
(DNA) and 201	/UG=Hs.28491 spermidinespermine N1-	
(amino acid)	acetyltransferase /FL=gb:M55580.1	
(amino acid)	acetyltransterase/FL-g0:M33380.1	

COT CAR 1.1 A	1 ND 6 0040 (0 1 /DDD II	201.420
COL6A3: alpha 3	gb:NM_004369.1 /DEF=Homo sapiens	201438_at
type VI collagen	collagen, type VI, alpha 3 (COL6A3), mRNA.	
isoform 1 precursor	/FEA=mRNA /GEN=COL6A3	
(LOC1293)	/PROD=collagen, type VI, alpha 3	
,	/DB XREF=gi:4758027 /UG=Hs.80988	
SEQ ID NOS: 55	collagen, type VI, alpha 3	
(DNA) and 202	/FL=gb:NM 004369.1	
(amino acid)	71L go.1\1\1_00+507.1	
	1 ND4 002 110 1 /DDF II	200665 = -4
SPARC: secreted	gb:NM_003 118.1 /DEF=Homo sapiens secreted	200665_s_at
protein, acidic,	protein, acidic, cysteine-rich (osteonectin)	
cysteine-rich	(SPARC), mRNA. /FEA=mRNA	
(osteonectin)	/GEN=SPARC /PROD=secreted protein, acidic,	
(LOC6678)	cysteine-rich(osteonectin)	
)	/DB XREF=gi:4507170 /UG=Hs.111779	
SEQ ID NOS: 56	secreted protein, acidic, cysteine-rich	
(DNA) and 203	(osteonectin) /FL=gb:BC004974.1 gb:J03040.1	
(amino acid)	gb:NM 003 118.1	
TXNIP: thioredoxin	gb:NM 006472.1 /DEF=Homo sapiens	201010 s at
	10 ~	201010_s_at
interacting protein	upregulated by 1,25-dihydroxyvitamin D-3	
(LOC10628)	(VDUP1), mRNA. /FEA=mRNA	
	/GEN=VDUP1 /PROD=upregulated by 1,25-	
SEQ ID NOS: 57	dihydroxyvitamin D-3 /DB_XREF=gi:5454161	
(DNA) and 204	/UG=Hs.179526 upregulated by 1,25-	
(amino acid)	dihydroxyvitamin D-3 /FL=gb:NM 006472.1	
,	gb:S73591.1	
MDK: midkine	gb:M69148.1 /DEF=Human midkine mRNA,	209035 at
(neurite growth-	complete cds. /FEA=mRNA /GEN=hMK-1	
promoting factor 2)	/PROD=midkine /DB XREF=gi:182650	
1 ~ /	/UG=Hs.82O45 midkine (neurite growth-	
(LOC4192)	`	
GTG T 110G 50	promoting factor 2) /FL=gb:M69148.1	
SEQ ID NOS: 58	gb:NM_002391.1	
(DNA) and 205		
(amino acid)		
TXNRD1:	gb:NM_003330.1 /DEF=Homo sapiens	201266_at
thioredoxin reductase	thioredoxin reductase 1 (TXNRD1), mRNA.	
1 (LOC7296)	/FEA=mRNA /GEN=TXNRD1	
- (=	/PROD=thioredoxin reductase 1	
SEQ ID NOS: 59	/DB XREF=gi:4507746 /UG=Hs.13046	
(DNA) and 206	thioredoxin reductase 1 /FL=gb:D88687.1	
1 ` ′	_	
(amino acid)	gb:AF077367.1 gb:NM_003330.1	
	gb:AF208018.1	200007
ARHD: ras homolog	gb:BC001338.1 /DEF=Homo sapiens, ras	209885_at
D (LOC29984)	homolog gene family, member, clone	
	MGC:5612, mRNA, complete cds.	
SEQ ID NOS: 60	/FEA=mRNA /PROD=ras homolog gene	
(DNA) and 207	family, member /DB_XREF=gi:12654980	
(amino acid)	/UG=Hs.15114 ras homolog gene family,	
	member /FL=gb:BC001338.1 gb:NM_014578.1	
	1110111001/1 13 60.D001330.1 60.1 1111_01 13/0.1	<u> </u>

PSPHL:	gb:NM 003832.1 /DEF=Homo sapiens	205048 s at
phosphoserine	phosphoserine phosphatase-like (PSPHL),	200010_5_4
phosphatase-like	mRNA. /FEA=mRNA /GEN=PSPHL	
(LOC8781)	/PROD=L-3-phosphoserine phosphatase	
(2000/01)	homolog /DB XREF=gi:4502934	
SEQ ID NOS: 61	/UG=Hs.76845 phosphoserine phosphatase-like	
(DNA) and 208	/FL=gb:NM 003832.1	
(amino acid)	/1 L-go.1010_003632.1	
RAB25: RAB25	chiNM 020297 1 /DEE-Home conions CATY	210106 of
	gb:NM_020387.1 /DEF=Homo sapiens CATX-	218186_at
(LOC57111)	8 protein (CATX-8), mRNA. /FEA=mRNA	
SEC ID NOS. 62	/GEN=CATX-8 /PROD=CATX-8 protein	
SEQ ID NOS: 62	/DB_XREF=gi:9966860 /UG=Hs.150826	
(DNA) and 209	CATX-8 protein /FL=gb:AF083124.1	
(amino acid)	gb:NM_020387.1	202026
SPINT1: hepatocyte	gb:NM_003710.1 /DEF=Homo sapiens serine	202826_at
growth factor	protease inhibitor, Kunitz type 1 (SPINT1),	
activator inhibitor 1	mRNA. /FEA=mRNA /GEN=SPINT1	
isoform 2 precursor	/PROD=hepatocyte growth factor activator	
(LOC6692)	inhibitorprecursor /DB_XREF=gi:4504328	
	/UG=Hs.233950 serine protease inhibitor,	
SEQ ID NOS: 63	Kunitz type 1 /FL=gb:BC004140.1	
(DNA) and 210	gb:AB000095.1 gb:NM_003710.1	
(amino acid)		
	1 170070071 7777	
SPINT2: serine	gb:AF027205.1 /DEF=Homo sapiens Kunitz-	210715_s_at
SPINT2: serine protease inhibitor,	gb:AF027205.1 /DEF=Homo sapiens Kunitz- type protease inhibitor (kop) mRNA, complete	210715_s_at
i	1 -	210715_s_at
protease inhibitor,	type protease inhibitor (kop) mRNA, complete	210715_s_at
protease inhibitor, Kunitz type, 2	type protease inhibitor (kop) mRNA, complete cds. /FEA=mRNA /GEN=kop /PROD=Kunitz-	210715_s_at
protease inhibitor, Kunitz type, 2	type protease inhibitor (kop) mRNA, complete cds. /FEA=mRNA /GEN=kop /PROD=Kunitz-type protease inhibitor /DB_XREF=gi:2598967	210715_s_at
protease inhibitor, Kunitz type, 2 (LOC10653)	type protease inhibitor (kop) mRNA, complete cds. /FEA=mRNA /GEN=kop /PROD=Kunitz-type protease inhibitor /DB_XREF=gi:2598967 /UG=Hs.31439 serine protease inhibitor, Kunitz	210715_s_at
protease inhibitor, Kunitz type, 2 (LOC10653) SEQ ID NOS: 64 (DNA) and 211	type protease inhibitor (kop) mRNA, complete cds. /FEA=mRNA /GEN=kop /PROD=Kunitz-type protease inhibitor /DB_XREF=gi:2598967 /UG=Hs.31439 serine protease inhibitor, Kunitz	210715_s_at
protease inhibitor, Kunitz type, 2 (LOC10653) SEQ ID NOS: 64 (DNA) and 211 (amino acid)	type protease inhibitor (kop) mRNA, complete cds. /FEA=mRNA /GEN=kop /PROD=Kunitz-type protease inhibitor /DB_XREF=gi:2598967 /UG=Hs.31439 serine protease inhibitor, Kunitz type, 2 /FL=gb:AF027205.1	
protease inhibitor, Kunitz type, 2 (LOC10653) SEQ ID NOS: 64 (DNA) and 211 (amino acid) EMP3: epithelial	type protease inhibitor (kop) mRNA, complete cds. /FEA=mRNA /GEN=kop /PROD=Kunitz-type protease inhibitor /DB_XREF=gi:2598967 /UG=Hs.31439 serine protease inhibitor, Kunitz type, 2 /FL=gb:AF027205.1 gb:NM_001425.1 /DEF=Homo sapiens	210715_s_at 203729_at
protease inhibitor, Kunitz type, 2 (LOC10653) SEQ ID NOS: 64 (DNA) and 211 (amino acid) EMP3: epithelial membrane protein 3	type protease inhibitor (kop) mRNA, complete cds. /FEA=mRNA /GEN=kop /PROD=Kunitz-type protease inhibitor /DB_XREF=gi:2598967 /UG=Hs.31439 serine protease inhibitor, Kunitz type, 2 /FL=gb:AF027205.1 gb:NM_001425.1 /DEF=Homo sapiens epithelial membrane protein 3 (EMP3), mRNA.	
protease inhibitor, Kunitz type, 2 (LOC10653) SEQ ID NOS: 64 (DNA) and 211 (amino acid) EMP3: epithelial	type protease inhibitor (kop) mRNA, complete cds. /FEA=mRNA /GEN=kop /PROD=Kunitz-type protease inhibitor /DB_XREF=gi:2598967 /UG=Hs.31439 serine protease inhibitor, Kunitz type, 2 /FL=gb:AF027205.1 gb:NM_001425.1 /DEF=Homo sapiens epithelial membrane protein 3 (EMP3), mRNA. /FEA=mRNA /GEN=EMP3 /PROD=epithelial	
protease inhibitor, Kunitz type, 2 (LOC10653) SEQ ID NOS: 64 (DNA) and 211 (amino acid) EMP3: epithelial membrane protein 3 (LOC2014)	type protease inhibitor (kop) mRNA, complete cds. /FEA=mRNA /GEN=kop /PROD=Kunitz-type protease inhibitor /DB_XREF=gi:2598967 /UG=Hs.31439 serine protease inhibitor, Kunitz type, 2 /FL=gb:AF027205.1 gb:NM_001425.1 /DEF=Homo sapiens epithelial membrane protein 3 (EMP3), mRNA. /FEA=mRNA /GEN=EMP3 /PROD=epithelial membrane protein 3 /DB_XREF=gi:4503562	
protease inhibitor, Kunitz type, 2 (LOC10653) SEQ ID NOS: 64 (DNA) and 211 (amino acid) EMP3: epithelial membrane protein 3 (LOC2014) SEQ ID NOS: 65	type protease inhibitor (kop) mRNA, complete cds. /FEA=mRNA /GEN=kop /PROD=Kunitz-type protease inhibitor /DB_XREF=gi:2598967 /UG=Hs.31439 serine protease inhibitor, Kunitz type, 2 /FL=gb:AF027205.1 gb:NM_001425.1 /DEF=Homo sapiens epithelial membrane protein 3 (EMP3), mRNA. /FEA=mRNA /GEN=EMP3 /PROD=epithelial membrane protein 3 /DB_XREF=gi:4503562 /UG=Hs.9999 epithelial membrane protein 3	
protease inhibitor, Kunitz type, 2 (LOC10653) SEQ ID NOS: 64 (DNA) and 211 (amino acid) EMP3: epithelial membrane protein 3 (LOC2014) SEQ ID NOS: 65 (DNA) and 212	type protease inhibitor (kop) mRNA, complete cds. /FEA=mRNA /GEN=kop /PROD=Kunitz-type protease inhibitor /DB_XREF=gi:2598967 /UG=Hs.31439 serine protease inhibitor, Kunitz type, 2 /FL=gb:AF027205.1 gb:NM_001425.1 /DEF=Homo sapiens epithelial membrane protein 3 (EMP3), mRNA. /FEA=mRNA /GEN=EMP3 /PROD=epithelial membrane protein 3 /DB_XREF=gi:4503562 /UG=Hs.9999 epithelial membrane protein 3 /FL=gb:U52101.1 gb:U87947.1	
protease inhibitor, Kunitz type, 2 (LOC10653) SEQ ID NOS: 64 (DNA) and 211 (amino acid) EMP3: epithelial membrane protein 3 (LOC2014) SEQ ID NOS: 65 (DNA) and 212 (amino acid)	type protease inhibitor (kop) mRNA, complete cds. /FEA=mRNA /GEN=kop /PROD=Kunitz-type protease inhibitor /DB_XREF=gi:2598967 /UG=Hs.31439 serine protease inhibitor, Kunitz type, 2 /FL=gb:AF027205.1 gb:NM_001425.1 /DEF=Homo sapiens epithelial membrane protein 3 (EMP3), mRNA. /FEA=mRNA /GEN=EMP3 /PROD=epithelial membrane protein 3 /DB_XREF=gi:4503562 /UG=Hs.9999 epithelial membrane protein 3 /FL=gb:U52101.1 gb:U87947.1 gb:NM_001425.1	203729_at
protease inhibitor, Kunitz type, 2 (LOC10653) SEQ ID NOS: 64 (DNA) and 211 (amino acid) EMP3: epithelial membrane protein 3 (LOC2014) SEQ ID NOS: 65 (DNA) and 212 (amino acid) TENS1: tensin-like	type protease inhibitor (kop) mRNA, complete cds. /FEA=mRNA /GEN=kop /PROD=Kunitz-type protease inhibitor /DB_XREF=gi:2598967 /UG=Hs.31439 serine protease inhibitor, Kunitz type, 2 /FL=gb:AF027205.1 gb:NM_001425.1 /DEF=Homo sapiens epithelial membrane protein 3 (EMP3), mRNA. /FEA=mRNA /GEN=EMP3 /PROD=epithelial membrane protein 3 /DB_XREF=gi:4503562 /UG=Hs.9999 epithelial membrane protein 3 /FL=gb:U52101.1 gb:U87947.1 gb:NM_001425.1 gb:NM_022748.1 /DEF=Homo sapiens	
protease inhibitor, Kunitz type, 2 (LOC10653) SEQ ID NOS: 64 (DNA) and 211 (amino acid) EMP3: epithelial membrane protein 3 (LOC2014) SEQ ID NOS: 65 (DNA) and 212 (amino acid) TENS1: tensin-like SH2 domain-	type protease inhibitor (kop) mRNA, complete cds. /FEA=mRNA /GEN=kop /PROD=Kunitz-type protease inhibitor /DB_XREF=gi:2598967 /UG=Hs.31439 serine protease inhibitor, Kunitz type, 2 /FL=gb:AF027205.1 gb:NM_001425.1 /DEF=Homo sapiens epithelial membrane protein 3 (EMP3), mRNA. /FEA=mRNA /GEN=EMP3 /PROD=epithelial membrane protein 3 /DB_XREF=gi:4503562 /UG=Hs.9999 epithelial membrane protein 3 /FL=gb:U52101.1 gb:U87947.1 gb:NM_001425.1 gb:NM_001425.1 gb:NM_01425.1 gb:NM_0142	203729_at
protease inhibitor, Kunitz type, 2 (LOC10653) SEQ ID NOS: 64 (DNA) and 211 (amino acid) EMP3: epithelial membrane protein 3 (LOC2014) SEQ ID NOS: 65 (DNA) and 212 (amino acid) TENS1: tensin-like SH2 domain- containing 1	type protease inhibitor (kop) mRNA, complete cds. /FEA=mRNA /GEN=kop /PROD=Kunitz-type protease inhibitor /DB_XREF=gi:2598967 /UG=Hs.31439 serine protease inhibitor, Kunitz type, 2 /FL=gb:AF027205.1 gb:NM_001425.1 /DEF=Homo sapiens epithelial membrane protein 3 (EMP3), mRNA. /FEA=mRNA /GEN=EMP3 /PROD=epithelial membrane protein 3 /DB_XREF=gi:4503562 /UG=Hs.9999 epithelial membrane protein 3 /FL=gb:U52101.1 gb:U87947.1 gb:NM_001425.1 gb:NM_022748.1 /DEF=Homo sapiens hypothetical protein FLJ13732 similar to tensin (FLJ13732), mRNA. /FEA=mRNA	203729_at
protease inhibitor, Kunitz type, 2 (LOC10653) SEQ ID NOS: 64 (DNA) and 211 (amino acid) EMP3: epithelial membrane protein 3 (LOC2014) SEQ ID NOS: 65 (DNA) and 212 (amino acid) TENS1: tensin-like SH2 domain-	type protease inhibitor (kop) mRNA, complete cds. /FEA=mRNA /GEN=kop /PROD=Kunitz-type protease inhibitor /DB_XREF=gi:2598967 /UG=Hs.31439 serine protease inhibitor, Kunitz type, 2 /FL=gb:AF027205.1 gb:NM_001425.1 /DEF=Homo sapiens epithelial membrane protein 3 (EMP3), mRNA. /FEA=mRNA /GEN=EMP3 /PROD=epithelial membrane protein 3 /DB_XREF=gi:4503562 /UG=Hs.9999 epithelial membrane protein 3 /FL=gb:U52101.1 gb:U87947.1 gb:NM_001425.1 gb:NM_022748.1 /DEF=Homo sapiens hypothetical protein FLJ13732 similar to tensin (FLJ13732), mRNA. /FEA=mRNA /GEN=FLJ13732 /PROD=hypothetical protein	203729_at
protease inhibitor, Kunitz type, 2 (LOC10653) SEQ ID NOS: 64 (DNA) and 211 (amino acid) EMP3: epithelial membrane protein 3 (LOC2014) SEQ ID NOS: 65 (DNA) and 212 (amino acid) TENS1: tensin-like SH2 domain- containing 1 (LOC64759)	type protease inhibitor (kop) mRNA, complete cds. /FEA=mRNA /GEN=kop /PROD=Kunitz-type protease inhibitor /DB_XREF=gi:2598967 /UG=Hs.31439 serine protease inhibitor, Kunitz type, 2 /FL=gb:AF027205.1 gb:NM_001425.1 /DEF=Homo sapiens epithelial membrane protein 3 (EMP3), mRNA. /FEA=mRNA /GEN=EMP3 /PROD=epithelial membrane protein 3 /DB_XREF=gi:4503562 /UG=Hs.9999 epithelial membrane protein 3 /FL=gb:U52101.1 gb:U87947.1 gb:NM_001425.1 gb:NM_022748.1 /DEF=Homo sapiens hypothetical protein FLJ13732 similar to tensin (FLJ13732), mRNA. /FEA=mRNA /GEN=FLJ13732 /PROD=hypothetical protein FLJ13732 similar to tensin	203729_at
protease inhibitor, Kunitz type, 2 (LOC10653) SEQ ID NOS: 64 (DNA) and 211 (amino acid) EMP3: epithelial membrane protein 3 (LOC2014) SEQ ID NOS: 65 (DNA) and 212 (amino acid) TENS1: tensin-like SH2 domain- containing 1 (LOC64759) SEQ ID NOS: 66	type protease inhibitor (kop) mRNA, complete cds. /FEA=mRNA /GEN=kop /PROD=Kunitz-type protease inhibitor /DB_XREF=gi:2598967 /UG=Hs.31439 serine protease inhibitor, Kunitz type, 2 /FL=gb:AF027205.1 gb:NM_001425.1 /DEF=Homo sapiens epithelial membrane protein 3 (EMP3), mRNA. /FEA=mRNA /GEN=EMP3 /PROD=epithelial membrane protein 3 /DB_XREF=gi:4503562 /UG=Hs.9999 epithelial membrane protein 3 /FL=gb:U52101.1 gb:U87947.1 gb:NM_001425.1 gb:NM_022748.1 /DEF=Homo sapiens hypothetical protein FLJ13732 similar to tensin (FLJ13732), mRNA. /FEA=mRNA /GEN=FLJ13732 /PROD=hypothetical protein FLJ13732 similar to tensin /DB_XREF=gi:12232408 /UG=Hs.12210	203729_at
protease inhibitor, Kunitz type, 2 (LOC10653) SEQ ID NOS: 64 (DNA) and 211 (amino acid) EMP3: epithelial membrane protein 3 (LOC2014) SEQ ID NOS: 65 (DNA) and 212 (amino acid) TENS1: tensin-like SH2 domain- containing 1 (LOC64759)	type protease inhibitor (kop) mRNA, complete cds. /FEA=mRNA /GEN=kop /PROD=Kunitz-type protease inhibitor /DB_XREF=gi:2598967 /UG=Hs.31439 serine protease inhibitor, Kunitz type, 2 /FL=gb:AF027205.1 gb:NM_001425.1 /DEF=Homo sapiens epithelial membrane protein 3 (EMP3), mRNA. /FEA=mRNA /GEN=EMP3 /PROD=epithelial membrane protein 3 /DB_XREF=gi:4503562 /UG=Hs.9999 epithelial membrane protein 3 /FL=gb:U52101.1 gb:U87947.1 gb:NM_001425.1 gb:NM_022748.1 /DEF=Homo sapiens hypothetical protein FLJ13732 similar to tensin (FLJ13732), mRNA. /FEA=mRNA /GEN=FLJ13732 /PROD=hypothetical protein FLJ13732 similar to tensin	203729_at

TTTP1 A . 1	1.ND 4.001520.1 /DEE. II	200000 /
HIF1A: hypoxia-	gb:NM_001530.1 /DEF=Homo sapiens	200989_at
inducible factor 1,	hypoxia-inducible factor 1, alpha subunit (basic	
alpha subunit isoform	helix-loop-helix transcription factor) (HIF1A),	
1 (LOC3091)	mRNA. /FEA=mRNA /GEN=HIF1A	
	/PROD=hypoxia-inducible factor 1, alpha	
SEQ ID NOS: 67	subunit (basichelix-loop-helix transcription	
(DNA) and 214	factor) /DB XREF=gi:4504384	
(amino acid)	/UG=Hs.197540 hypoxia-inducible factor 1,	
	alpha subunit (basic helix-loop-helix	
	transcription factor) /FL=gb:U29165.1	
	gb:AF304431.1 gb:NM_001530.1	
	gb:AF207601.1 gb:AF207602.1 gb:U22431.1	
ST14: matriptase	gb:NM 021978.1 /DEF=Homo sapiens	202005 at
(LOC6768)	suppression of tumorigenicity 14 (colon	_
(2000,00)	carcinoma, matriptase, epithin) (ST14), mRNA.	
SEQ ID NOS: 68	/FEA=mRNA /GEN=ST14	
(DNA) and 215	/PROD=suppression of tumorigenicity 14	
(amino acid)	(coloncarcinoma, matriptase, epithin)	
	/DB_XREF=gi:11415039 /UG=Hs.56937	
	suppression of tumorigenicity 14 (colon	
	carcinoma, matriptase, epithin)	
	/FL=gb:AF057145.1 gb:NM 021978.1	
	gb:AB030036.1 gb:AF133086.1	
	gb:AF118224.2	
STK17A:	Consensus includes gb:AW194730 /FEA=EST	202693 s at
serine/threonine	/DB_XREF=gi:6473630	202075_5_at
kinase 17a	/DB_XREF=est:xn43d11.x1	
(apoptosis-inducing)	/CLONE=IMAGE:2696469 /UG=Hs.9075	
(LOC9263)	serinethreonine kinase 17a (apoptosis-inducing)	
	/FL=gb:AB011420.1 gb:NM_004760.1	
SEQ ID NOS: 69		
(DNA) and 216		
(amino acid)		
SH3YL1:	gb:NM 015677.1 /DEF=Homo sapiens	204019 s at
hypothetical protein	hypothetical protein (DKFZP586F1318),	
DKFZP586F1318	mRNA./FEA=mRNA	
(LOC26751)	/GEN=DKFZP586F1318 /PROD=hypothetical	
GEO TO TYPE TO	protein /DB_XREF=gi:7661669 /UG=Hs.25213	
SEQ ID NOS: 70	hypothetical protein /FL=gb:NM_015677.1	
(DNA) and 217		
(amino acid)		
EXT1: exostoses	gb:NM 000127.1 /DEF=Homo sapiens	201995_at
(multiple) 1	exostoses (multiple) 1 (EXT1), mRNA.	_
(LOC2131)	/FEA=mRNA /GEN=EXT1 /PROD=exostoses	
	(multiple) 1 /DB XREF=gi·4557570	
	(multiple) 1 /DB_XREF=gi:4557570	
SEQ ID NOS: 71	/UG=Hs.184161 exostoses (multiple) 1	
	`	

	1 NR 6 017 102 1 /DEE II	010010
GALNT7:	gb:NM_017423.1 /DEF=Homo sapiens UDP-	218313_s_at
polypeptide N-	N-acetyl-alpha-D-galactosamine:polypeptide	
acetylgalactosaminylt	N-acetylgalactosaminyltransferase 7 (GalNAc-	
ransferase 7	T7) (GALNT7), mRNA. /FEA=mRNA	
(LOC51809)	/GEN=GALNT7/PROD=polypeptide N-	
(E0031007)	acetylgalactosaminyltransferase 7	
GEO 777 3 10 G 70	,	
SEQ ID NOS: 72	/DB_XREF=gi:8393408 /UG=Hs.246315 UDP-	
(DNA) and 219	N-acetyl-alpha-D-galactosamine:polypeptide	
(amino acid)	N-acetylgalactosaminyltransferase 7 (GalNAc-	
	T7) /FL=gb:NM 017423.1	
SDC1: syndecan 1	gb:NM 002997.1 /DEF=Homo sapiens	201287 s at
(LOC6382)	syndecan 1 (SDC1), mRNA. /FEA=mRNA	
(LOCO382)	/GEN=SDC1 /PROD=syndecan 1	
GEO 770 3 10 G 770		
SEQ IID NOS: 73	/DB_XREF=gi:4506858 /UG=Hs.82109	
(DNA) and 220	syndecan 1 /FL=gb:J05392.1 gb:NM_002997.1	
(amino acid)		
ITGAV: integrin,	Consensus includes gb:AI093579 /FEA=EST	202351 at
alpha V (vitronectin	/DB XREF=gi:3432555	_
receptor, alpha	/DB XREF=est:qb15g06.x1	
	/CLONE=IMAGE:1696378/UG=Hs.295726	
polypeptide, antigen	* · - · · · ·	
CD51) (LOC3685)	integrin, alpha V (vitronectin receptor, alpha	
	polypeptide, antigen CD51) /FL=gb:M14648.1	
SEQ IID NOS: 74	gb:NM 002210.1	
(DNA) and 221		
(amino acid)		
ANXA6: annexin VI	gb:NM 001155.2 /DEF=Homo sapiens annexin	200982 s at
	10 -	200962_s_at
isoform 1 (LOC309)	A6 (ANXA6), transcript variant 1, mRNA.	,
	/FEA=mRNA /GEN=ANXA6 /PROD=annexin	
SEQ ID NOS: 75	VI isoform 1 /DB_XREF=gi:4809274	
(DNA) and 222	/UG=Hs.118796 annexin A6 /FL=gb:J03578.1	
(amino acid)	gb:D00510.1 gb:NM_001155.2	
PDGF C: platelet-	gb:NM 016205.1 /DEF=Homo sapiens platelet	218718 at
derived growth factor	derived growth factor C (PDGFC), mRNA.	
	/FEA=mRNA /GEN=PDGFC	
C precursor		
(LOC56034)	/PROD=secretory growth factor-like protein	
	fallotein /DB_XREF=gi:9994186	
SEQ ID NOS: 76	/UG=Hs.43080 platelet derived growth factor C	
(DNA) and 223	/FL=gb:AF091434.1 gb:AF244813.1	
(amino acid)	gb:AB033831.1 gb:NM 016205.1	
FLNA: filamin 1	Consensus includes gb:AI625550 /FEA=EST	214752 x at
•		211132_A_at
(actin-binding	/DB_XREF=gi:4650481	
protein-280)	/DB_XREF=est:ty57d06.x1	
(LOC2316)	/CLONE=IMAGE:2283179 /UG=Hs.195464	
	filamin A, alpha (actin-binding protein-280)	
SEQ ID NOS: 77	_ ` ` ` ` ` ` ` ` ` ` ` ` ` ` ` ` ` ` `	
		1
(DNA) and 224		
(DNA) and 224 (amino acid)		

FLNA: filamin 1 (actin-binding protein-280) (LOC2316) SEQ ID NOS: 78 (DNA) and 225 (amino acid)	Consensus includes gb:AW051856 /FEA=EST /DB_XREF=gi:5914215 /DB_XREF=est:wz04a05.x1 /CLONE=IMAGE:2557040 /UG=Hs.195464 filamin A, alpha (actin-binding protein-280)	213746_s_at
TUBA3: tubulin, alpha 3 (LOC7846) SEQ ID NOS: 79 (DNA) and 226 (amino acid)	gb:AF141347.1 /DEF=Homo sapiens hum-a- tub2 alpha-tubulin mRNA, complete cds. /FEA=mRNA /PROD=alpha-tubulin /DB_XREF=gi:4929133 /UG=Hs.272897 Tubulin, alpha, brain-specific /FL=gb:AF141347.1 gb:NM_006009.1	209118_s_at
LOXL2: lysyl oxidase-like 2 (LOC4017) SEQ ID NOS: 80 (DNA) and 227 (amino acid)	gb:NM_002318.1 /DEF=Homo sapiens lysyl oxidase-like 2 (LOXL2), mRNA. /FEA=mRNA /GEN=LOXL2 /PROD=lysyl oxidase-like 2 /DB_XREF=gi:4505010 /UG=Hs.83354 lysyl oxidase-like 2 /FL=gb:BC000594.1 gb:U89942.1 gb:NM_002318.1 gb:AF1 17949.1	202998_s_at
CYR61: cysteinerich, angiogenic inducer, 61 (LOC3491) SEQ ID NOS: 81 (DNA) and 228 (amino acid)	gb:AF003114.1 /DEF=Homo sapiens CYR61 mRNA, complete cds. /FEA=mRNA /GEN=CYR61 /DB_XREF=gi:6649848 /UG=Hs.8867 cysteine-rich, angiogenic inducer, 61 /FL=gb:AF003114.1	210764_s_at
GALNT3: polypeptide N- acetylgalactosaminylt ransferase 3 (LOC2591) SEQ ID NOS: 82 (DNA) and 229 (amino acid)	Consensus includes gb:BF063271 /FEA=EST /DB_XREF=gi:10822181 /DB_XREF=est:7h87d05.x1 /CLONE=IMAGE:3322953 /UG=Hs.278611 UDP-N-acetyl-alpha-D-galactosamine:polypeptide N-acetylgalactosaminyltransferase 3 (GalNAc-T3) /FL=gb:NM_004482.2	203397_s_at
MAP1B: microtubule- associated protein 1B isoform 1 (LOC4131) SEQ ID NOS: 83 (DNA) and 230 (amino acid)	Consensus includes gb:AL523076 /FEA=EST /DB_XREF=gi:12786569 /DB_XREF=est:AL523076 /CLONE=CS0DC001YI12 (3 prime) /UG=Hs.82503 H.sapiens mRNA for 3UTR of unknown protein	212233_at

TUBB-5: tubulin	gb:BC002654.1 /DEF=Homo sapiens, Similar	209191 at
beta-5 (LOC84617)	to tubulin, beta, 4, clone MGC:4083, mRNA,	200101_00
	complete cds. /FEA=mRNA /PROD=Similar to	
SEQ ID NOS: 84	tubulin, beta, 4 /DB XREF=gi:12803638	
(DNA) and 231	/UG=Hs.274398 Homo sapiens, Similar to	
(amino acid)	tubulin, beta, 4, clone MGC:4083, mRNA,	
(animo acid)	complete cds /FL=gb:BC002654.1	
TYMS: thymidylate	gb:NM 001071.1 /DEF=Homo sapiens	202589 at
synthetase	thymidylate synthetase (TYMS), mRNA.	202309_at
(LOC7298)	/FEA=mRNA /GEN=TYMS	
(LOC/298)	/PROD=thymidylate synthetase	
SEO ID NIOS: 95	/PROD-utylindylate synthetase /DB XREF=gi:4507750 /UG=Hs.82962	
SEQ ID NOS: 85		
(DNA) and 232	thymidylate synthetase /FL=gb:BC002567.1	'
(amino acid)	gb:NM_001071.1	00.6000
IFI16: interferon,	gb:NM_005531.1 /DEF=Homo sapiens	206332_s_at
gamma-inducible	interferon, gamma-inducible protein 16 (IFI16),	
protein 16	mRNA. /FEA=mRNA /GEN=IFI16	
(LOC3428)	/PROD=interferon, gamma-inducible protein 16	
	/DB_XREF=gi:5031778 /UG=Hs.155530	
SEQ ID NOS: 86	interferon, gamma-inducible protein 16	
(DNA) and 233	/FL=gb:M63838.1 gb:NM_005531.1	
(amino acid)		
GRB10: growth	gb:D86962.1 /DEF=Human mRNA for	209409_at
factor receptor-bound	KIAA0207 gene, complete cds. /FEA=mRNA	
protein 1O	/GEN=KIAA0207 /DB_XREF=gi:1503997	
(LOC2887)	/UG=Hs.81875 growth factor receptor-bound	
	protein 10 /FL=gb:D86962.1 gb:AF000017.1	'
SEQ ID NOS: 87		
(DNA) and 234		
(amino acid)		
FLNA: filamin 1	gb:NM_001456.1 /DEF=Homo sapiens filamin	200859_x_at
(actin-binding	A, alpha (actin-binding protein-280) (FLNA),	
protein-280)	mRNA. /FEA=mRNA /GEN=FLNA	
(LOC2316)	/PROD=filamin 1 (actin-binding protein-280)	
	/DB XREF=gi:4503744 /UG=Hs.195464	
SEQ ID NOS: 88	filamin A, alpha (actin-binding protein-280)	
(DNA) and 235	/FL=gb:NM 001456.1	j.
(amino acid)		
TNC: tenascin C	gb:NM 002160.1 /DEF=Homo sapiens	201645 at
(hexabrachion)	hexabrachion (tenascin C, cytotactin) (HXB),	
(LOC3371)	mRNA. /FEA=mRNA /GEN=HXB	
	/PROD=hexabrachion (tenascin C, cytotactin)	
SEQ ID NOS: 89	/DB_XREF=gi:4504548 /UG=Hs.289114	
(DNA) and 236	hexabrachion (tenascin C, cytotactin)	
(amino acid)	/FL=gb:M55618.1 gb:NM_002160.1	
(millio acia)	/1 D 50.11133010.1 50.1 1111_002100.1	L

	<u></u>	
SLC26A2: sulfate	Consensus includes gb:AI025519 /FEA=EST	205097_at
anion transporter 1	/DB_XREF=gi:3241132	
(LOC1836)	/DB_XREF=est:ov75c04.x1	
	/CLONE=IMAGE:1643142 /UG=Hs.29981	
SEQ ID NOS: 90	solute carrier family 26 (sulfate transporter),	
(DNA) and 237	member 2 /FL=gb:NM_000112.1 gb:U14528.1	
(amino acid)		
KIAA0 7 46:	Consensus includes gb:AB018289.1	212314_at
KIAA0746 protein	/DEF=Homo sapiens mRNA for KIAA0746	
(LOC23 231)	protein, partial cds. /FEA=mRNA	
	/GEN=KIAA0746 /PROD=KIAA0746 protein	
SEQ ID NOS: 91	/DB_XREF=gi:3882212 /UG=Hs.49500	
(DNA) and 238	KIAA0746 protein	
(amino acid)	_	
LAMP1: lysosomal-	gb:NM_005561.2 /DEF=Homo sapiens	201553_s_at
associated membrane	lysosomal-associated membrane protein 1	
protein 1 (LOC3916)	(LAMP1), mRNA. /FEA=mRNA	
	/GEN=LAMP1 /PROD=lysosomal-associated	
SEQ ID NOS: 92	membrane protein 1 /DB XREF=gi:7669500	
(DNA) and 239	/UG=Hs.150101 lysosomal-associated	
(amino acid)	membrane protein 1 /FL=gb:J04182.1	
	gb:J03263.1 gb:NM_005561.2	
DPYSL2:	gb:NM_001386.1 /DEF=Homo sapiens	200762_at
dihydropyrimidinase-	dihydropyrimidinase-like 2 (DPYSL2), mRNA.	
like 2 (LOC1808)	/FEA=mRNA /GEN=DPYSL2	
	/PROD=dihydropyrimidinase-like 2	
SEQ ID NOS: 93	/DB_XREF=gi:4503376 /UG=Hs.173381	
(DNA) and 240	dihydropyrimidinase-like 2 /FL=gb:U17279.1	
(amino acid)	gb:D78013.1 gb:U97105.1 gb:NM_001386.1	
IFI16: interferon,	gb:AF208043.1 /DEF=Homo sapiens IFI16b	208966_x_at
gamma-inducible	(IFI16b) mRNA, complete cds. /FEA=mRNA	
protein 16	/GEN=IFI16b/PROD=IFI16b	
(LOC3428)	/DB_XREF=gi:6644296 /UG=Hs.155530	
	interferon, gamma-inducible protein 16	
SEQ ID NOS: 94	/FL=gb:AF208043.1	
(DNA) and 241		
(amino acid)		
KPNB2: karyopherin	Consensus includes gb:AI307759 /FEA=EST	221829_s_at
beta 2 (LOC3842)	/DB_XREF=gi:4002363	
	/DB_XREF=est:tb24g08.x1	
SEQ ID NOS: 95	/CLONE=IMAGE:2055326 /UG=Hs.168075	
(DNA) and 242	karyopherin (importin) beta 2	
(amino acid)		:

DDND.	1. ATA 4. 000211 1 /DEE 11	201200
PRNP: prion protein	gb:NM_000311.1 /DEF=Homo sapiens prion	20130O_s_at
preproprotein	protein (p27-30) (Creutzfeld-Jakob disease,	
(LOC5621)	Gerstmann-Strausler-Scheinker syndrome, fatal	:
	familial insomnia) (PRNP), mRNA.	
SEQ ID NOS: 96	/FEA=mRNA /GEN=PRNP /PROD=prion	
(DNA) and 243	protein /DB_XREF=gi:4506112 /UG=Hs.74621	
(amino acid)	prion protein (p27-30) (Creutzfeld-Jakob	
	disease, Gerstmann-Strausler-Scheinker	
	syndrome, fatal familial insomnia)	
	/FL=gb:AY008282.1 gb:M13899.1	
	gb:NM_000311.1	
RAI14: retinoic acid	gb:NM_015577.1 /DEF=Homo sapiens novel	202052_s_at
induced 14	retinal pigment epithelial gene (NORPEG),	
(LOC26064)	mRNA. /FEA=mRNA /GEN=NORPEG	
	/PROD=DKFZP564G013 protein	
SEQ ID NOS: 97	/DB XREF=gi:13470085 /UG=Hs.15165 novel	
(DNA) and 244	retinal pigment epithelial gene	1
(amino acid)	/FL=gb:NM 015577.1 gb:AF155135.1	
JAG1: jagged 1	gb:U61276.1 /DEF=Human transmembrane	209098 s at
precursor (LOC182)	protein Jagged 1 (HJ1) mRNA, complete cds.	Service Students
	/FEA=mRNA/GEN=HJ1	
SEQ ID NOS: 98	/PROD=transmembrane protein Jagged 1	
(DNA) and 245	/DB XREF=gi:1438936 /UG=Hs.91143 jagged	
(amino acid)	1 (Alagille syndrome) /FL=gb:U61276.1	
(daring dela)	gb:U73936.1 gb:AF003837.1 gb:AF028593.1	
	gb:NM 000214.1	
CLIC4: chloride	gb:NM 013943.1 /DEF=Homo sapiens chloride	20156O at
intracellular channel	intracellular channel 4 (CLIC4), mRNA.	201300_at
4 (LOC25932)	/FEA=mRNA /GEN=CLIC4 /PROD=chloride	
(LOC23932)	intracellular channel 4 /DB XREF=gi:7330334	
SEQ ID NOS: 99	/UG=Hs.25035 chloride intracellular channel 4	
(DNA) and 246	/FL=gb:AF109196.1 gb:AF097330.1	
(amino acid)	gb:AL117424.1 gb:NM_013943.1	
TP53I3: tumor	gb:BC000474.1 /DEF=Homo sapiens, quinone	210609 s at
protein p53 inducible		210009_s_ai
protein 3 (LOC9540)	oxidoreductase homolog, clone MGC:8642, mRNA, complete cds. /FEA=mRNA	j.
protein 3 (LOC9340)	/PROD=quinone oxidoreductase homolog	
SEQ ID NOS: 100	/PROD—quinone oxidoreductase nomolog /DB XREF=gi:12653408 /UG=Hs.50649	
(DNA) and 247	quinone oxidoreductase homolog	
1 '	, <u> </u>	
(amino acid) EFA6R: ADP-	/FL=gb:BC000474.1 Consensus includes gb:AW117368 /FEA=EST	202254 - c+
	1	203354_s_at
ribosylation factor	/DB_XREF=gi:6085952	
guanine nucleotide	/DB_XREF=est:xd88h01.x1	
factor 6 (LOC23362)	/CLONE=IMAGE:2604721 /UG=Hs.6763	
GEO ID NIOS: 101	KIAA 0942 protein /FL=gb:AF243495.2	
SEQ ID NOS: 101	gb:NM_015310.1	
(DNA) and 248		
(amino acid)		

JUP: junction	gb:NM 02199 1.1 /DEF=Homo sapiens junction	201015 s at
plakoglobin	plakoglobin (JUP), transcript variant 2, mRNA.	201015_5_6
(LOC3728)	/FEA=mRNA /GEN=JUP /PROD=junction	
(2003/20)	plakoglobin, is oform 1	
SEQ ID NOS: 102	/DB XREF=g i :12056467 /UG=Hs.2340	
(DNA) and 249	junction plakoglobin /FL=gb:NM_021991.1	
(amino acid)	gb:BC000441. 1	
PAPSS2: 3'-	gb:NM 00467 0.1 /DEF=Homo sapiens 3-	203059 s at
		203039_S_att
phosphoadenosine 5'-	phosphoadenosine 5-phosphosulfate synthase 2	
phosphosulfate	(PAPSS2), mRNA. /FEA=mRNA	
synthase 2	/GEN=PAPSS2 /PROD=3-prime-	
(LOC9060)	phosphoadenosine 5-prime-	
GEO TO MOG. 100	phosphosulfatesynthase 2	
SEQ ID NOS: 103	/DB_XREF=gi:4758879 /UG=Hs.274230 3-	
(DNA) and 250	phosphoadenosine 5-phosphosulfate synthase 2	
(amino acid)	/FL=gb:AF15O754.2 gb:AF313907.1	
	gb:AF091242. 1 gb:NM_004670.1	
	gb:AF074331. 1 gb:AF173365.1	
DKK3: dickkopf	Consensus includes gb:AU148057 /FEA=EST	214247_s_at
homolog 3	/DB_XREF=g i :11009578	
(LOC27122)	/DB_XREF=est:AU148057	
	/CLONE=MAMMA1002489 /UG=Hs.278503	
SEQ ID NOS: 104	regulated in glioma	
(DNA) and 251		
(amino acid)		
JAG1: jagged 1	Consensus includes gb:U77914.1 /DEF=Human	216268_s_at
precursor (LOC182)	soluble protein Jagged mRNA, partial cds.	
	/FEA=mRNA /PROD=soluble protein Jagged	
SEQ ID NOS: 105	/DB XREF=gi:1684889/UG=Hs.91143 jagged	
(DNA) and 252	1 (Alagille syndrome)	
(amino acid)		
CALD1: caldesmon 1	Consensus includes gb:AL583520 /FEA=EST	212077 at
isoform 3 (LOC800)	/DB XREF=g i :12952562	_
	/DB XREF=est:AL583520	
SEQ ID NOS: 106	/CLONE=CS0IDC024YE13 (5 prime)	
(DNA) and 253	/UG=Hs.182183 Homo sapiens mRNA for	
(amino acid)	caldesmon, 3 UTR	
DPYSL3:	Consensus includes gb:W72516 /FEA=EST	201430 s at
dihydropyrimidinase-	/DB XREF=gi:1382173	
like 3 (LOC1809)	/DB XREF=est:zd64g05.s1	
(200100)	/CLONE=IMA GE:345464 /UG=Hs.74566	
SEQ ID NOS: 107	dihydropyrimidinase-like 3 /FL=gb:D78014.1	
(DNA) and 254	gb:NM_001387.1	
(amino acid)	50.11111_00130 7.1	
(ammo acid)	<u></u>	

PMP22: peripheral myelin protein 22 (LOC5376) SEQ ID NOS: 108 (DNA) and 255 (amino acid)	gb:L03203.1 /DEF=Human peripheral myelin protein 22 (G.AS3) mRNA, complete cds. /FEA=mRNA /GEN=GAS3 /PROD=peripheral myelin protein 22 /DB_XREF=gi:182984 /UG=Hs.103724 peripheral myelin protein 22 /FL=gb:L032 03.1	210139_s_at
ALCAM: activated leukocyte cell adhesion molecule (LOC214) SEQ ID NOS: 109 (DNA) and 256 (amino acid)	Consensus includes gb:BF242905 /FEA=EST /DB_XREF=gi:11156833 /DB_XREF=est:601877949F1 /CLONE=IM_AGE:4106028 /UG=Hs.10247 activated leucocyte cell adhesion molecule /FL=gb:NM_001627.1 gb:L38608.1	201951_at
PAPSS2: 3'- phosphoadenosine 5'- phosphosulfate synthase 2 (LOC9060) SEQ ID NOS: 110 (DNA) and 257 (amino acid)	Consensus includes gb:AW299958 /FEA=EST /DB_XREF=gi:6709635 /DB_XREF=est:xs44g05.x1 /CLONE=IM_AGE:2772536 /UG=Hs.274230 3-phosphoadenosine 5-phosphosulfate synthase 2 /FL=gb:AF150754.2 gb:AF313907.1 gb:AF091242.1 gb:NM_004670.1 gb:AF074331.1 gb:AF173365.1	203058_s_at
KPNB2: karyopherin beta 2 (LOC3842) SEQ ID NOS: 111 (DNA) and 258 (amino acid)	gb:NM_0022 70.1 /DEF=Homo sapiens karyopherin (importin) beta 2 (KPNB2), mRNA. /FEA=mRNA /GEN=KPNB2 /PROD=karyopherin (importin) beta 2 /DB_XREF=gi:4504906 /UG=Hs.168075 karyopherin (importin) beta 2 /FL=gb:U703 22.1 gb:NM_002270.1	207657_x_at
PTPRE: protein tyrosine phosphatase, receptor type, E isoform 1 precursor (LOC5791) SEQ ID NOS: 112 (DNA) and 259	Consensus in cludes gb:AA775177 /FEA=EST /DB_XREF=gi:2834511 /DB_XREF=est:ac79a06.s1 /CLONE=IM_AGE:868786 /UG=Hs.31137 protein tyrosine phosphatase, receptor type, E /FL=gb:NM_006504.1	221840_at
(amino acid) TRB2: tribbles homolog 2 (LOC28951) SEQ ID NOS: 113 (DNA) and 260 (amino acid)	gb:NM_021643.1 /DEF=Homo sapiens GS3955 protein (GS3955), mRNA. /FEA=mRNA /GEN=GS3955 /PROD=GS3955 protein /DB_XREF=gi:11056053 /UG=Hs.155418 GS3955 protein /FL=gb:NM_021643.1 gb:BC002637.1 gb:D87119.1	202478_at

COL13A1: alpha 1	gb:M33653.1 /DEF=Human (clones HT-	211343 s at
type XIII collagen	125,133) alpha-2 type IV collagen (COL4A2)	211515_5_4
isoform 1 (LOC130)	mRNA, complete cds. /FEA=mRNA	
isololiii i (isociso)	/GEN=COL4A2 /PROD=alpha-2 type IV	·
SEQ ID NOS: 114	collagen /DB XREF=gi:180828	
(DNA) and 261	/UG=Hs.211933 collagen, type XIII, alpha 1	
(amino acid)	/FL=gb:M33653.1	
		202760 s at
PALM2: paralemmin	gb:NM_007203.1 /DEF=Homo sapiens A	202760_s_at
2 (LOC114299)	kinase (PRKA) anchor protein 2 (AKAP2), mRNA. /FEA=mRNA /GEN=AKAP2	
GEO TO MOS. 115		
SEQ ID NOS: 115	/PROD=A kinase (PRKA) anchor protein 2	
(DNA) and 262	/DB_XREF=gi:6005708 /UG=Hs.42322 A	
(amino acid)	kinase (PRKA) anchor protein 2	
	/FL=gb:AB023137.1 gb:NM_007203.1	001667
GJA1: connexin 43	gb:NM_000165.2 /DEF=Homo sapiens gap	201667_at
(LOC2697)	junction protein, alpha 1, 43kD (connexin 43)	
	(GJA1), mRNA. /FEA=mRNA/GEN=GJA1	
SEQ ID NOS: 116	/PROD=connexin 43 /DB_XREF=gi:4755136	
(DNA) and 263	/UG=Hs.74471 gap junction protein, alpha 1,	
(amino acid)	43kD (connexin 43) /FL=gb:M65188.1	
	gb:NM_000165.2	
FLJ10901:	gb:NM_018265.1 /DEF=Homo sapiens	219010_at
hypothetical protein	hypothetical protein FLJ1O901 (FLJ10901),	
FLJ10901	mRNA./FEA=mRNA/GEN=FLJ10901	
(LOC55765)	/PROD=hypothetical protein FLJ10901	
, l	/DB_XREF=gi:8922753 /UG=Hs.73239	
SEQ ID NOS: 117	hypothetical protein FLJ1O901	
(DNA) and 264	/FL=gb:NM_018265.1	
(amino acid)	,	
EFEMP1: EGF-	Consensus includes gb:AI826799 /FEA=EST	201842_s_at
containing fibulin-	/DB_XREF=gi:5447470	
like extracellular	$DB_XREF=est:wk56d07.x1$	
matrix protein 1	/CLONE=IMAGE:24194 0 5 /UG=Hs.76224	
isoform a precursor	EGF-containing fibulin-like extracellular matrix	
(LOC2202)	protein 1 /FL=gb:U03877. 1 gb:NM_004105.2	
SEQ ID NOS: 118		
(DNA) and 265		
(amino acid)		
NRP1: neuropilin 1	Consensus includes gb:BE 620457 /FEA=EST	212298_at
(LOC8829)	/DB_XREF=gi:9891395	
	/DB_XREF=est:60148369 0F1	
SEQ ID NOS: 119	/CLONE=IMAGE:388605 5 /UG=Hs.69285	
(DNA) and 266	neuropilin 1 /FL=gb:AF01 8956.1	
(amino acid)	gb:AF016050.1 gb:NM 0O3873.1	

CLDN7: claudin 7	gb:NM 001307.1 /DEF=Honno sapiens claudin	202790 at
(LOC1366)	7 (CLDN7), mRNA. /FEA=n1RNA	
(2001300)	/GEN=CLDN7/PROD=claudin 7	
SEQ ID NOS: 120	/DB XREF=gi:10835007/UG=Hs.278562	
(DNA) and 267	claudin 7 /FL=gb:NM 001307.1	
(amino acid)	gb:BC001055.1	
CED-6: PTB domain	gb:NM 016315.1 /DEF=Homo sapiens CED-6	204237 at
adaptor protein CED-	protein (CED-6), mRNA. /FEA=mRNA	20-1257_at
6 (LOC51454)	/GEN=CED-6/PROD=CED-6 protein	
0 (EOC31434)	/DB XREF=gi:7705317 /UG=Hs.107056 CED-	
SEQ ID NOS: 121	6 protein /FL=gb:AF200715. 1 gb:AF191771.1	
(DNA) and 268	gb:NM 016315.1	
(amino acid)	gb.1NN_010313.1	
CSPG2: chondroitin	Consensus includes gb:BF59O263 /FEA=EST	204619 s at
	/DB XREF=gi:11682587	204019_s_at
sulfate proteoglycan 2 (versican) (LOC1462)	/DB_XREF=g1.11082387 /DB_XREF=est:nab22b12.x1	
(versicali) (LOC1402)	/CLONE=IMAGE:3266638/UG=Hs.81800	
SEQ ID NOS: 122		
1 -	chondroitin sulfate proteogly can 2 (versican)	
(DNA) and 269	/FL=gb:NM_004385.1	
(amino acid)	-1-1172060 1 /DEE-Hyman Lagrandaria bata?	200226 a at
KPNB2: karyopherin	gb:U72069.1 /DEF=Human karyopherin beta2	209226_s_at
beta 2 (LOC3842)	mRNA, complete cds. /FEA==mRNA	
GEO ID NICE 102	/PROD=karyopherin beta2	
SEQ ID NOS: 123	/DB_XREF=gi:1657775 /UG=Hs.168075	
(DNA) and 270	karyopherin (importin) beta 2	
(amino acid)	/FL=gb:U72069.1 gb:U72395.1	010717
MLAT4: myxoid	gb:NM_018192.1 /DEF=Horno sapiens	218717_s_at
liposarcoma	hypothetical protein FLJ1071 8 (FLJ10718),	
associated protein 4	mRNA. /FEA=mRNA /GEN=FLJ10718	
(LOC55214)	/PROD=hypothetical protein FLJ10718	
	/DB_XREF=gi:8922618 /UG=Hs.42824	
SEQ ID NOS: 124	hypothetical protein FLJ1071 8	
(DNA) and 271	/FL=gb:NM_018192.1	
(amino acid)	1 30 1505 1 / DDD 11	010006
TPM1: tropomyosin 1	gb:Z24727.1 /DEF=H.sapien.s tropomyosin	210986_s_at
(alpha) (LOC7168)	isoform mRNA, complete CDS. /FEA=mRNA	
GEO ID NICE 127	/PROD=tropomyosin isoform	
SEQ ID NOS: 125	/DB_XREF=gi:854188 /UG=Hs.77899	
(DNA) and 272	tropomyosin 1 (alpha) /FL=g b :Z24727.1	
(amino acid)	1 375 0170(4.1 /7)777 1 1 377 0	206504
LY96: MD-2 protein	gb:NM_015364.1 /DEF=Horno sapiens MD-2	206584_at
(LOC23643)	protein (MD-2), mRNA. /FEA=mRNA	
CTO TT 3.700	/GEN=MD-2 /PROD=MD-2 protein	
SEQ ID NOS: 126	/DB_XREF=gi:7662503 /UG=Hs.69328 MD-2	
(DNA) and 273	protein /FL=gb:AB018549.1 gb:NM_015364.1	
(amino acid)	gb:AF168121.1	

COL6A1: collagen,	Consensus includes gb:AI141603 / FEA=EST	212091 s at
type VI, alpha 1	/DB XREF=gi:3649060	
precursor (LOC1291)	/DB XREF=est:qa90h10.x1	
	/CLONE=IMAGE:1694083 /UG= I Hs.108885	
SEQ ID NOS: 127	collagen, type VI, alpha 1	
(DNA) and 274		
(amino acid)		
CDC42EP3: Cdc42	gb:AL136842.1 /DEF=Homo sapiens mRNA;	209288 s at
effector protein 3	cDNA DKFZp434A0530 (from clone	
(LOC10602)	DKFZp434A0530); complete cds.	
	/FEA=mRNA /GEN=DKFZp434A0530	
SEQ ID NOS: 128	/PROD=hypothetical protein	
(DNA) and 275	/DB XREF=gi:6807668 /UG=Hs,260024	
(amino acid)	Cdc42 effector protein 3 /FL=gb:AF094521.1	
	gb:AF104857.1 gb:NM 006449.1	
	gb:AF164118.1 gb:AL136842.1	
JTB: jumping	gb:NM 006694.1 /DEF=Homo sapiens	200048 s at
translocation	jumping translocation breakpoint (JTB),	
breakpoint	mRNA. /FEA=mRNA /GEN=JTB	
(LOC10899)	/PROD=jumping translocation breakpoint	
	/DB_XREF=gi:5729888 /UG=Hs.6396 jumping	
SEQ ID NOS: 129	translocation breakpoint /FL=gb:B C000499.1	
(DNA) and 276	gb:BC001363.1 gb:BC000996.2	
(amino acid)	gb:BC001667.1 gb:AB016488.1	
	gb:AF131797.1 gb:NM_006694.1	
	gb:AF115850.2	
CDH2: cadherin 2,	gb:M34064.1 /DEF=Human N-cad herin	203440_at
type 1 preproprotein	mRNA, complete cds. /FEA=mRNA	
(LOC1000)	/GEN=NCAD /DB_XREF=gi:416292	
	/UG=Hs.161 cadherin 2, type 1, N-cadherin	
SEQ ID NOS: 130	(neuronal) /FL=gb:M34064.1 gb:NM_001792.1	
(DNA) and 277		
(amino acid)		
MYLK: myosin light	gb:NM_005965.1 /DEF=Homo sapiens myosin,	202555_s_at
chain kinase isoform	light polypeptide kinase (MYLK), mRNA.	
6 (LOC4638)	/FEA=mRNA /GEN=MYLK /PROD=myosin,	
	light polypeptide kinase	
SEQ ID NOS: 131	/DB_XREF=gi:5174600 /UG=Hs.211582	
(DNA) and 278	myosin, light polypeptide kinase	
(amino acid)	/FL=gb:AB037663.1 gb:NM_005965.1	
	gb:AF069601.2	

COL4A1: alpha 1	Consensus includes gb:NM_001845.1	2 1 1981 at
type IV collagen	/DEF=Homo sapiens collagen, type IV, alpha 1	
preproprotein	(COL4A1), mRNA. /FEA=CDS	
(LOC1282)	/GEN=COL4A1 /PROD=collagen, type IV,	
(2001202)	alpha 1 /DB_XREF=gi:7656984	
SEQ ID NOS: 132	/UG=Hs.119129 collagen, type IV, alpha 1	
(DNA) and 279	/FL=gb:NM 001845.1	
(amino acid)	712 goului_0010 1011	
PROS1: protein S	gb:NM 000313.1 /DEF=Homo sapiens protein	2O7808 s at
(alpha) (LOC5627)	S (alpha) (PROS1), mRNA. /FEA=mRNA	20,000_5_00
(шрш) (2000-1)	/GEN=PROS1 /PROD=protein S (alpha)	
SEQ ID NOS: 133	/DB XREF=gi:4506116 /UG=Hs.64016 protein	
(DNA) and 280	S (alpha) /FL=gb:M15036.1 gb:NM 000313.1	
(amino acid)	S (mpm) /1 2 goint to our goin in z_ocourson	
EFEMP1: EGF-	gb:NM_004105.2 /DEF=Homo sapiens EGF-	2O1843 s at
containing fibulin-	containing fibulin-like extracellular matrix	
like extracellular	protein 1 (EFEMP1), transcript variant 1,	
matrix protein 1	mRNA. /FEA=mRNA /GEN=EFEMP1	
isoform a precursor	/PROD=EGF-containing fibulin-like	
(LOC2202)	extracellular matrixprotein 1 precursor, isoform	
	a precursor /DB_XREF=gi:9665261	
SEQ ID NOS: 134	/UG=Hs.76224 EGF-containing fibulin-like	
(DNA) and 281	extracellular matrix protein 1 /FL=gb:U03877.1	
(amino acid)	gb:NM 004105.2	
CCL2: small	Consensus includes gb:S69738.1 /DEF=MCP-	2 1 6598 s at
inducible cytokine A2	1=monocyte chemotactic protein human, aortic	
precursor (LOC6347)	endothelial cells, mRNA, 661 nt. /FEA=mRNA	
	/GEN=MCP-1 /PROD=MCP-1	
SEQ ID NOS: 135	/DB_XREF=gi:545464 /UG=Hs.303649 small	
(DNA) and 282	inducible cytokine A2 (monocyte chemotactic	
(amino acid)	protein 1, homologous to mouse Sig-je)	
DFNA5: deafness,	gb:NM_004403.1 /DEF=Homo sapiens	2O3695_s_at
autosomal dominant 5	deafness, autosomal dominant 5 (DFNA5),	
protein (LOC1687)	mRNA. /FEA=mRNA /GEN=DFNA5	
	/PROD=deafness, autosomal dominant 5	
SEQ ID NOS: 136	protein /DB_XREF=gi:4758153 /UG=Hs.13530	
(DNA) and 283	deafness, autosomal dominant 5	
(amino acid)	/FL=gb:AF073308.1 gb:NM_004403.1	
-	gb:AF007790.2	
TPM1: tropomyosin 1	gb:M19267.1 /DEF=Human tropomyosin	2 1 0987_x_at
(alpha) (LOC7168)	mRNA, complete cds. /FEA=mRNA	
	/DB_XREF=gi:339943 /UG=Hs.77899	
SEQ ID NOS: 137	tropomyosin 1 (alpha) /FL=gb:M19267.1	
(DNA) and 284		
(amino acid)		

DDAH1:	Consensus includes gb:AL078459	209094_at
dimethylarginine	/DEF=Human DNA sequence from clone RP4-	
dimethylaminohydrol	621F18 on chromosome 1p11.4-21.3. Contains	
ase 1 (LOC23576)	the 3 end of the gene for ng,ng	'
	dimethylarginine dimethylaminohydrolase (EC	
SEQ ID NOS: 138	3.5.3.18), ESTs, STSs and GSSs /FEA=mRNA	
(DNA) and 285	/DB XREF=gi:5791502 /UG=Hs.303180	
(amino acid)	dimethylarginine dimethylaminohydrolase 1	
()	/FL=gb:AB001915.1 gb:NM 012137.1	
PMAIP1: phorbol-12-	Consensus includes gb:AI857639 /FEA=EST	204285 s at
myristate-13-acetate-	/DB XREF=gi:5511255	201203_5_4
induced protein 1	/DB_XREF=est:wk95g09.x1	
	/CLONE=IMAGE:2423200 /UG=Hs.96	
(LOC5366)		
GEO ID NOG 120	phorbol-12-myristate-13-acetate-induced	
SEQ ID NOS: 139	protein 1 /FL=gb:NM_021127.1	
(DNA) and 286		
(amino acid)		
ACOX2: acyl-	gb:NM_003500.1 /DEF=Homo sapiens acyl-	205364_at
Coenzyme A oxidase	Coenzyme A oxidase 2, branched chain	
2, branched chain	(ACOX2), mRNA. /FEA=mRNA	:
(LOC8309)	/GEN=ACOX2 /PROD=acyl-Coenzyme A	
	oxidase 2, branched chain	
SEQ ID NOS: 140	/DB XREF=gi:4501868 /UG=Hs.9795 acyl-	
(DNA) and 287	Coenzyme A oxidase 2, branched chain	
(amino acid)	/FL=gb:NM 003500.1	
GDI1: GDP	gb:NM 001493.1 /DEF=Homo sapiens GDP	201864 at
dissociation inhibitor	dissociation inhibitor 1 (GDI1), mRNA.	_
1 (LOC2664)	/FEA=mRNA /GEN=GDI1 /PROD=GDP	
1 (2002001)	dissociation inhibitor 1 /DB XREF=gi:4503970	
SEQ ID NOS: 141	/UG=Hs.74576 GDP dissociation inhibitor 1	
(DNA) and 288	/FL=gb:BC000317.1 gb:NM_001493.1	
(amino acid)	gb:D45021.1	
DPYSL3:	gb:NM 001387.1 /DEF=Homo sapiens	201431 s at
1	dihydropyrimidinase-like 3 (DPYSL3), mRNA.	201431_8_at
dihydropyrimidinase-	, , , , , , , , , , , , , , , , , , , ,	
like 3 (LOC1809)	/FEA=mRNA /GEN=DPYSL3	
GEO ID NOC 140	/PROD=dihydropyrimidinase-like 3	
SEQ ID NOS: 142	/DB_XREF=gi:4503378 /UG=Hs.74566	
(DNA) and 289	dihydropyrimidinase-like 3 /FL=gb:D78014.1	
(amino acid)	gb:NM_001387.1	010550
APOC1:	Consensus includes gb:W79394 /FEA=EST	213553_x_at
apolipoprotein C-I	/DB_XREF=gi:1390665	
precursor (LOC341)	/DB_XREF=est:zd80c07.s1	
	/CLONE=IMAGE:346956 /UG=Hs.268571	
SEQ ID NOS: 143	apolipoprotein C-I	
(DNA) and 290		
(amino acid)		
· · · · · · · · · · · · · · · · · · ·		

TTC3:	gb:NM_003316.1 /DEF=Homo sapiens	208073_x_aet
tetratricopeptide	tetratricopeptide repeat domain 3 (TTC3),	
repeat domain 3	mRNA. /FEA=mRNA /GEN=TTC3	
(LOC7267)	/PROD=tetratricopeptide repeat domain 3	
	/DB XREF=gi:10835036 /UG=Hs.118174	
SEQ ID NOS: 144	tetratricopeptide repeat domain 3	
(DNA) and 291	/FL=gb:NM 003316.1 gb:D84295.1	
(amino acid)		
SNX6: sorting nexin	gb:NM_021249.1 /DEF=Homo sapiens sorting	217789 at
6 isoform a	nexin 6 (SNX6), mRNA. /FEA=mRNA	
(LOC58533)	/GEN=SNX6 /PROD=sorting nexin 6	
	/DB XREF=gi:13027619 /UG=Hs.284291	
SEQ ID NOS: 145	sorting nexin 6 /FL=gb:BC001798.1	,
(DNA) and 292	gb:NM_021249.1 gb:AF121856.1	
(amino acid)		
CKAP4:	Consensus includes gb:AW029619 /FEA=EST	200998 s at
transmembrane	/DB XREF=gi:5888375	
protein (63kD),	/DB_XREF=est:wx14e05.x1	
endoplasmic	/CLONE=IMAGE:2543648 /UG=Hs.74368	
reticulum/Golgi	transmembrane protein (63kD), endoplasmic	
interm (LOC10970)	reticulumGolgi intermediate compartment	
	/FL=gb:NM_006825.1	
SEQ ID NOS: 146		
(DNA) and 293		
(amino acid)		
TUBB: tubulin, beta	gb:NM_001069.1 /DEF=Homo sapiens tubulin,	204141_at
polypeptide	beta polypeptide (TUBB), mRNA.	
(LOC7280)	/FEA=mRNA /GEN=TUBB /PROD=tubulin,	
	beta polypeptide /DB_XREF=gi:4507728	
SEQ ID NOS: 147	/UG=Hs.179661 tubulin, beta polypeptide	
(DNA) and 294	/FL=gb:BC001194.1 gb:NM_001069.1	
(amino acid)		

The biomarkers provided in Table 1, which include the nucleotide sequences of SEQ ID NOS:1-147 and the amino acid sequences of SEQ ID NOS:148-294, referred to herein as a total of 147 biomarkers with reference to the Unigene Title, includes 40 cases where multiple probe sets measure the intensity of a single biomarker (at most, three probe sets for one biomarker). In these cases, the redundant probe sets reference the same full-length cDNA and protein sequences. Table 2 provides a correlation between the NCBI locus IDs and the probe set IDs.

TABLE 2 - Correlation between NCBI Locus IDs and Probe Set IDs

NCBI	Number of	Probe set IDs
	Probe sets	11000 500 125
182	3	209099 x at, 209098 s at, 216268 s at
1462	3	204620 s at, 221731 x at, 204619 s at
2316	3	214752 x at, 213746 s at, 200859 x at
3842	3	221829 s at, 207657 x at, 209226 s at
9060	3	203060 s at, 203059 s at, 203058 s at
10899	3	210434_x_at, 210927_x_at, 200048_s_at
214	2	201952_at, 201951_at
1291	2	213428_s_at, 212091_s_at
1508	2	200839_s_at, 200838_at
1809	2	201430_s_at, 201431_s_at
2202	2	201842_s_at, 201843_s_at
2810	2	33322_i_at, 33323_r_at
3428	2	206332_s_at, 208966_x_at
3491	2	201289_at, 210764_s_at
7168	2	210986_s_at, 210987_x_at
8781	2	212509_s_at, 205048_s_at
10628	2	201008_s_at, 201010_s_at
130	1	211343_s_at
309	1	200982_s_at
341	1	213553_x_at
754	1	200677_at
800	1	212077_at
999	1	201131_s_at
1000	1	203440_at
1075	1	201487_at
1282	1	211981_at
1292	1	209156_s_at
1293	1	201438_at
1366	1	202790_at
1490	1	209101_at
1687	11	203695_s_at
1808	1	200762_at
1836	1	205097_at
2014	1	203729 at
2115	1	221911_at
2131	11	201995_at
2150	1	213506 at
2273	1	201540_at
2591	1	203397 s at
2664	1	201864_at
2697	1	201667_at
2791	1	204115_at
2887	1	209409_at

3091	1	200989 at
3371	1	201645 at
3383	1	202637 s at
3576	1	211506 s at
3685	1	202351_at
3728	1	201015_s_at
3855	1	209016_s_at
3856	1	209008_x_at
3875	1	201596_x_at
3880	1	201650_at
3916	1	201553_s_at
4017	1	202998_s_at
4131	1	212233_at
4192	1	209035_at
4638	1	202555_s_at
5066	1	202336_s_at
5270	1	212190_at
5366	1	204285 s at
5376	1	210139 s at
5441	1	211730 s at
5621	$\overline{1}$	201300 s at
5627	1	207808 s at
5791	1	221840 at
5834	1	201481 s at
6137	1	212191 x at
6159	1	213969 x at
6280	1	203535 at
6303	1	210592 s at
6347	1	216598 s at
6382	1	201287 s at
6535		
	1	202219 at
6678	1	200665 s at
6692		202826_at
6748	<u>l</u>	201004_at
6768	1	202005_at
6772	1	200887_s_at
7045	1	201506_at
7267	1	208073_x_at
7280	1	204141_at
7296	1	201266_at
7298	1	202589_at
7345	111	201387 s at
7358	1	203343_at
7846	1	209118_s_at
8309	1_	205364_at
8829	1	212298_at
9263	1	202693 s at

9540	1	210609_s_at
10135	1	217738_at
10602	1	209288_s_at
10653	1	210715_s_at
10962	1	211071_s_at
10970	1	200998_s_at
11098	1	202458_at
11199	1	210143_at
22943	1	204602_at
23231	1	212314_at
23362	1	203354_s_at
23576	1	209094_at
23643	1	206584_at
25932	1	201560_at
26064	1	202052_s_at
26751	1	204019_s_at
27122	1	214247_s_at
28951	1	202478_at
29984	1	209885_at
51065	1	218007_s_at
51454	1	204237_at
51809	1	218313_s_at
54407	1	218041_x_at
55214	1	218717_s_at
55765	1	219010_at
56034	1	218718_at
57111	1	218186_at
57402	1	218677_at
58533	1	217789_at
64759	1	217853_at
84617	1	209191_at
113146	1	212992_at
114299	1	202760_s_at
347902	1	222108_at

The biomarkers have expression levels in the cells that may be dependent on the activity of the EGFR signal transduction pathway, and that are also highly correlated with EGFR modulator sensitivity exhibited by the cells. Biomarkers serve as useful molecular tools for predicting a response to EGFR modulators, preferably biological molecules, small molecules, and the like that affect EGFR kinase activity via direct or indirect inhibition or antagonism of EGFR kinase function or activity.

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EGFR MODULATORS

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As used herein, the term "EGFR modulator" is intended to mean a compound or drug that is a biological molecule or a small molecule that directly or indirectly modulates EGFR activity or the EGFR signal transduction pathway. Thus, compounds or drugs as used herein is intended to include both small molecules and biological molecules. Direct or indirect modulation includes activation or inhibition of EGFR activity or the EGFR signal transduction pathway. In one aspect, inhibition refers to inhibition of the binding of EGFR to an EGFR ligand such as, for example, EGF. In another aspect, inhibition refers to inhibition of the kinase activity of EGFR.

EGFR modulators include, for example, EGFR-specific ligands, small molecule EGFR inhibitors, and EGFR monoclonal antibodies. In one aspect, the EGFR modulator inhibits EGFR activity and/or inhibits the EGFR signal transduction pathway. In another aspect, the EGFR modulator is an EGFR monoclonal antibody that inhibits EGFR activity and/or inhibits the EGFR signal transduction pathway.

EGFR modulators include biological molecules or small molecules. Biological molecules include all lipids and polymers of monosaccharides, amino acids, and nucleotides having a molecular weight greater than 450. Thus, biological molecules include, for example, oligosaccharides and polysaccharides; oligopeptides, polypeptides, peptides, and proteins; and oligonucleotides and polynucleotides. Oligonucleotides and polynucleotides include, for example, DNA and RNA.

Biological molecules further include derivatives of any of the molecules described above. For example, derivatives of biological molecules include lipid and glycosylation derivatives of oligopeptides, polypeptides, peptides, and proteins.

Derivatives of biological molecules further include lipid derivatives of oligosaccharides and polysaccharides, e.g., lipopolysaccharides. Most typically, biological molecules are antibodies, or functional equivalents of antibodies. Functional equivalents of antibodies have binding characteristics comparable to those of antibodies, and inhibit the growth of cells that express EGFR. Such functional equivalents include, for example, chimerized, humanized, and single chain antibodies as well as fragments thereof.

Functional equivalents of antibodies also include polypeptides with amino acid sequences substantially the same as the amino acid sequence of the variable or

hypervariable regions of the antibodies. An amino acid sequence that is substantially the same as another sequence, but that differs from the other sequence by means of one or more substitutions, additions, and/or deletions, is considered to be an equivalent sequence. Preferably, less than 50%, more preferably less than 25%, and still more preferably less than 10%, of the number of amino acid residues in a sequence are substituted for, added to, or deleted from the protein.

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The functional equivalent of an antibody is preferably a chimerized or humanized antibody. A chimerized antibody comprises the variable region of a non-human antibody and the constant region of a human antibody. A humanized antibody comprises the hypervariable region (CDRs) of a non-human antibody. The variable region other than the hypervariable region, e.g., the framework variable region, and the constant region of a humanized antibody are those of a human antibody.

Suitable variable and hypervariable regions of non-human antibodies may be derived from antibodies produced by any non-human mammal in which monoclonal antibodies are made. Suitable examples of mammals other than humans include, for example, rabbits, rats, mice, horses, goats, or primates.

Functional equivalents further include fragments of antibodies that have binding characteristics that are the same as, or are comparable to, those of the whole antibody. Suitable fragments of the antibody include any fragment that comprises a sufficient portion of the hypervariable (i.e., complementarity determining) region to bind specifically, and with sufficient affinity, to EGFR tyrosine kinase to inhibit growth of cells that express such receptors.

Such fragments may, for example, contain one or both Fab fragments or the $F(ab')_2$ fragment. Preferably, the antibody fragments contain all six complementarity determining regions of the whole antibody, although functional fragments containing fewer than all of such regions, such as three, four, or five CDRs, are also included.

In one aspect, the fragments are single chain antibodies, or Fv fragments. Single chain antibodies are polypeptides that comprise at least the variable region of the heavy chain of the antibody linked to the variable region of the light chain, with or without an interconnecting linker. Thus, Fv fragment comprises the entire antibody combining site. These chains may be produced in bacteria or in eukaryotic cells.

The antibodies and functional equivalents may be members of any class of immunoglobulins, such as IgG, IgM, IgA, IgD, or IgE, and the subclasses thereof. In one aspect, the antibodies are members of the IgG1 subclass. The functional equivalents may also be equivalents of combinations of any of the above classes and subclasses.

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In one aspect, EGFR antibodies can be selected from chimerized, humanized, fully human, and single chain antibodies derived from the murine antibody 225 described in U.S. Patent No. 4,943,533 to Mendelsohn et al.

In another aspect, the EGFR antibody can be selected from the antibodies described in U.S. Patent No. 6,235,883 to Jakobovits et al., U.S. Patent No. 5,558,864 to Bendi et al., and U.S. Patent No. 5,891,996 to Mateo de Acosta del Rio et al.

In addition to the biological molecules discussed above, the EGFR modulators useful in the invention may also be small molecules. Any molecule that is not a biological molecule is considered herein to be a small molecule. Some examples of small molecules include organic compounds, organometallic compounds, salts of organic and organometallic compounds, saccharides, amino acids, and nucleotides. Small molecules further include molecules that would otherwise be considered biological molecules, except their molecular weight is not greater than 450. Thus, small molecules may be lipids, oligosaccharides, oligopeptides, and oligonucleotides and their derivatives, having a molecular weight of 450 or less.

It is emphasized that small molecules can have any molecular weight. They are merely called small molecules because they typically have molecular weights less than 450. Small molecules include compounds that are found in nature as well as synthetic compounds. In one embodiment, the EGFR modulator is a small molecule that inhibits the growth of tumor cells that express EGFR. In another embodiment, the EGFR modulator is a small molecule that inhibits the growth of refractory tumor cells that express EGFR.

Numerous small molecules have been described as being useful to inhibit EGFR. For example, U.S. Patent No. 5,656,655 to Spada et al. discloses styryl substituted heteroaryl compounds that inhibit EGFR. The heteroaryl group is a monocyclic ring with one or two heteroatoms, or a bicyclic ring with 1 to about 4 heteroatoms, the compound being optionally substituted or polysubstituted.

U.S. Patent No. 5,646,153 to Spada et al. discloses bis mono and/or bicyclic aryl heteroaryl, carbocyclic, and heterocarbocyclic compounds that inhibit EGFR.

- U.S. Patent No. 5,679,683 to Bridges et al. discloses tricyclic pyrimidine compounds that inhibit the EGFR. The compounds are fused heterocyclic pyrimidine derivatives described at column 3, line 35 to column 5, line 6.
- U.S. Patent No. 5,616,582 to Barker discloses quinazoline derivatives that have receptor tyrosine kinase inhibitory activity.

Fry et al., Science 265, 1093-1095 (1994) in Figure 1 discloses a compound having a structure that inhibits EGFR.

Osherov et al. disclose tyrphostins that inhibit EGFR/HER1 and HER 2, particularly those in Tables I, II, III, and IV.

U.S. Patent No. 5,196,446 to Levitzki et al. discloses heteroarylethenediyl or heteroarylethendeiylaryl compounds that inhibit EGFR, particularly from column 2, line 42 to column 3, line 40.

Panek et al., Journal of Pharmacology and Experimental Therapeutics 283, 1433-1444 (1997) discloses a compound identified as PD166285 that inhibits the EGFR, PDGFR, and FGFR families of receptors. PD166285 is identified as 6-(2,6-dichlorophenyl)-2-(4-(2-diethylaminoethyoxy)phenylamino)-8-methyl-8H-pyrido(2,3-d)pyrimidin-7-one having the structure shown in Figure 1 on page 1436.

BIOMARKERS AND BIOMARKER SETS

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The invention includes individual biomarkers and biomarker sets having both diagnostic and prognostic value in disease areas in which signaling through EGFR or the EGFR pathway is of importance, e.g., in cancers or tumors, in immunological disorders, conditions or dysfunctions, or in disease states in which cell signaling and/or cellular proliferation controls are abnormal or aberrant. The biomarker sets comprise a plurality of biomarkers such as, for example, a plurality of the biomarkers provided in Table 1, that highly correlate with resistance or sensitivity to one or more EGFR modulators.

The biomarker sets of the invention enable one to predict or reasonably foretell the likely effect of one or more EGFR modulators in different biological systems or for cellular responses. The biomarker sets can be used in *in vitro* assays of

EGFR modulator response by test cells to predict *in vivo* outcome. In accordance with the invention, the various biomarker sets described herein, or the combination of these biomarker sets with other biomarkers or markers, can be used, for example, to predict how patients with cancer might respond to therapeutic intervention with one or more EGFR modulators.

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A biomarker set of cellular gene expression patterns correlating with sensitivity or resistance of cells following exposure of the cells to one or more EGFR modulators provides a useful tool for screening one or more tumor samples before treatment with the EGFR modulator. The screening allows a prediction of cells of a tumor sample exposed to one or more EGFR modulators, based on the expression results of the biomarker set, as to whether or not the tumor, and hence a patient harboring the tumor, will or will not respond to treatment with the EGFR modulator.

The biomarker or biomarker set can also be used as described herein for monitoring the progress of disease treatment or therapy in those patients undergoing treatment for a disease involving an EGFR modulator.

The biomarkers also serve as targets for the development of therapies for disease treatment. Such targets may be particularly applicable to treatment of lung disease, such as non-small cell lung cancers or tumors. Indeed, because these biomarkers are differentially expressed in sensitive and resistant cells, their expression patterns are correlated with relative intrinsic sensitivity of cells to treatment with EGFR modulators. Accordingly, the biomarkers highly expressed in resistant cells may serve as targets for the development of new therapies for the tumors which are resistant to EGFR modulators, particularly EGFR inhibitors.

The level of biomarker protein and/or mRNA can be determined using methods well known to those skilled in the art. For example, quantification of protein can be carried out using methods such as ELISA, 2-dimensional SDS PAGE, Western blot, immunopreciptation, immunohistochemistry, fluorescence activated cell sorting (FACS), or flow cytometry. Quantification of mRNA can be carried out using methods such as PCR, array hybridization, Northern blot, in-situ hybridization, dot-blot, Taqman, or RNAse protection assay.

MICROARRAYS

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The invention also includes specialized microarrays, e.g., oligonucleotide microarrays or cDNA microarrays, comprising one or more biomarkers, showing expression profiles that correlate with either sensitivity or resistance to one or more EGFR modulators. Such microarrays can be employed in in vitro assays for assessing the expression level of the biomarkers in the test cells from tumor biopsies, and determining whether these test cells are likely to be resistant or sensitive to EGFR modulators. For example, a specialized microarray can be prepared using all the biomarkers, or subsets thereof, as described herein and shown in Table 1. Cells from a tissue or organ biopsy can be isolated and exposed to one or more of the EGFR modulators. Following application of nucleic acids isolated from both untreated and treated cells to one or more of the specialized microarrays, the pattern of gene expression of the tested cells can be determined and compared with that of the biomarker pattern from the control panel of cells used to create the biomarker set on the microarray. Based upon the gene expression pattern results from the cells that underwent testing, it can be determined if the cells show a resistant or a sensitive profile of gene expression. Whether or not the tested cells from a tissue or organ biopsy will respond to one or more of the EGFR modulators and the course of treatment or therapy can then be determined or evaluated based on the information gleaned from the results of the specialized microarray analysis.

ANTIBODIES

The invention also includes antibodies, including polyclonal or monoclonal, directed against one or more of the polypeptide biomarkers. Such antibodies can be used in a variety of ways, for example, to purify, detect, and target the biomarkers of the invention, including both *in vitro* and *in vivo* diagnostic, detection, screening, and/or therapeutic methods.

KITS

The invention also includes kits for determining or predicting whether a patient would be susceptible or resistant to a treatment that comprises one or more EGFR modulators. The patient may have a cancer or tumor such as, for example, a

non-small cell lung cancer or tumor. Such kits would be useful in a clinical setting for use in testing a patient's biopsied tumor or other cancer samples, for example, to determine or predict if the patient's tumor or cancer will be resistant or sensitive to a given treatment or therapy with an EGFR modulator. The kit comprises a suitable container that comprises: one or more microarrays, e.g., oligonucleotide microarrays or cDNA microarrays, that comprise those biomarkers that correlate with resistance and sensitivity to EGFR modulators, particularly EGFR inhibitors; one or more EGFR modulators for use in testing cells from patient tissue specimens or patient samples; and instructions for use. In addition, kits contemplated by the invention can further include, for example, reagents or materials for monitoring the expression of biomarkers of the invention at the level of mRNA or protein, using other techniques and systems practiced in the art such as, for example, RT-PCR assays, which employ primers designed on the basis of one or more of the biomarkers described herein, immunoassays, such as enzyme linked immunosorbent assays (ELISAs), immunoblotting, e.g., Western blots, or in situ hybridization, and the like, as further described herein.

APPLICATION OF BIOMARKERS AND BIOMARKER SETS

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The biomarkers and biomarker sets may be used in different applications. Biomarker sets can be built from any combination of biomarkers listed in Table 1 to make predictions about the likely effect of any EGFR modulator in different biological systems. The various biomarkers and biomarkers sets described herein can be used, for example, as diagnostic or prognostic indicators in disease management, to predict how patients with cancer might respond to therapeutic intervention with compounds that modulate the EGFR, and to predict how patients might respond to therapeutic intervention that modulates signaling through the entire EGFR regulatory pathway.

The biomarkers have both diagnostic and prognostic value in diseases areas in which signaling through EGFR or the EGFR pathway is of importance, e.g., in immunology, or in cancers or tumors in which cell signaling and/or proliferation controls have gone awry.

In accordance with the invention, cells from a patient tissue sample, e.g., a tumor or cancer biopsy, can be assayed to determine the expression pattern of one or more biomarkers prior to treatment with one or more EGFR modulators. In one aspect, the tumor or cancer is NSCLC. Success or failure of a treatment can be determined based on the biomarker expression pattern of the cells from the test tissue (test cells), e.g., tumor or cancer biopsy, as being relatively similar or different from the expression pattern of a control set of the one or more biomarkers. Thus, if the test cells show a biomarker expression profile which corresponds to that of the biomarkers in the control panel of cells which are sensitive to the EGFR modulator, it is highly likely or predicted that the individual's cancer or tumor will respond favorably to treatment with the EGFR modulator. By contrast, if the test cells show a biomarker expression pattern corresponding to that of the biomarkers of the control panel of cells which are resistant to the EGFR modulator, it is highly likely or predicted that the individual's cancer or tumor will not respond to treatment with the EGFR modulator.

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The invention also provides a method of monitoring the treatment of a patient having a disease treatable by one or more EGFR modulators. The isolated test cells from the patient's tissue sample, e.g., a tumor biopsy or tumor sample, can be assayed to determine the expression pattern of one or more biomarkers before and after exposure to an EGFR modulator wherein, preferably, the EGFR modulator is an EGFR inhibitor. The resulting biomarker expression profile of the test cells before and after treatment is compared with that of one or more biomarkers as described and shown herein to be highly expressed in the control panel of cells that are either resistant or sensitive to an EGFR modulator. Thus, if a patient's response is sensitive to treatment by an EGFR modulator, based on correlation of the expression profile of the one or biomarkers, the patient's treatment prognosis can be qualified as favorable and treatment can continue. Also, if, after treatment with an EGFR modulator, the test cells don't show a change in the biomarker expression profile corresponding to the control panel of cells that are sensitive to the EGFR modulator, it can serve as an indicator that the current treatment should be modified, changed, or even discontinued. This monitoring process can indicate success or failure of a patient's treatment with an EGFR modulator and such monitoring processes can be repeated as necessary or desired.

The biomarkers of the invention can be used to predict an outcome prior to having any knowledge about a biological system. Essentially, a biomarker can be considered to be a statistical tool. Biomarkers are useful primarily in predicting the phenotype that is used to classify the biological system.

Although the complete function of all of the biomarkers are not currently known, some of the biomarkers are likely to be directly or indirectly involved in the EGFR signaling pathway. In addition, some of the biomarkers may function in metabolic or other resistance pathways specific to the EGFR modulators tested. Notwithstanding, knowledge about the function of the biomarkers is not a requisite for determining the accuracy of a biomarker according to the practice of the invention.

EXAMPLES:

EXAMPLE 1 - Identification of Biomarkers

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The biomarkers of Table 1 were identified using three particular approaches. The transcriptional profiling data from primary tumors and cell lines was examined to identify genes with expression that is highly variable across the tumors and cell lines. In addition, attempts were made to determine the IC₅₀ on a panel of cell lines in order to identify genes whose expression profiles correlate with sensitive/resistant classification based on IC₅₀ values. Furthermore, cell lines and xenograft models were treated with the chimeric EGFR antibody cetuximab (marketed as Erbitux®) and the small molecule EGFR inhibitor gefitinib to identify genes that are modulated by EGFR inhibitors.

NSCLC tumors and patients:

RNAs from twenty-nine NSCLC adenocarcinoma tumors were obtained (Ardais Corporation, Somerville, MA). Adenocarcinomas are the most common subtype of NSCLC. The median age of the patients was 65 years (range: 43-80 years). The tumors belonged to all size ranges T1 – T4 and all stages ranging from Stage IA to Stage IV according to the AJCC classification.

Determination of Relative Drug Sensitivity in NSCLC Cell Lines:

The NSCLC cell lines were grown using standard cell culture conditions:

DMEM supplemented to contain 10% fetal bovine serum, 100 IU/ml penicillin, 100 mg/ml streptomycin and 2 mM L-glutamine (all from Invitrogen Life Technologies, Carlsbad, CA). Fourteen non-small cell lung cancer cell lines were examined for their sensitivity to EGFR inhibitor monoclonal antibody cetuximab. Cytotoxicity was assessed in cells by BrdU Cell Proliferation colorimetric ELISA (Roche Applied Science, Indianapolis, IN). This is a colorimetric immunoassay for the quantification of cell proliferation based on the measurement of BrdU incorporation during DNA synthesis. To carry out the assays, the NSCLC cells were plated at 2500-5000 cells/well in 96 well microtiter plates and 24 hours later diluted monoclonal antibody drug was added. The concentrations for the EGFR inhibitor cetuximab used in the cytotoxicity assays was 5 µg/ml, 4 µg/ml, 2 µg/ml, 1 µg/ml and 0.5 µg/ml. The cells were incubated at 37 °C for 48 hours at which time the BrdU labeling reagent was added. After two hours the labeling medium was removed and cells were fixed and the DNA was denatured using a FixDenat solution. The anti-BrdU antibody conjugated with peroxidase was added and immune complexes were detected by the subsequent substrate reaction. The reaction product was quantified by measuring the absorbance of the samples in an ELISA reader at 450 nm. The greater the absorbency, the greater the number of live cells. Only two of the fourteen cell lines tested had an IC₅₀ between 4 and $5\mu g/ml$. The IC₅₀ is the drug concentration required to inhibit cell proliferation to 50% of that of untreated cells. Three to six independent BrdU assays were performed for each cell line.

Resistance/sensitivity classification:

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FIG. 1 shows the mRNA level of the epidermal growth factor receptor gene as determined by expression profiling of fourteen NSCLC cell lines that were tested in the BrdU assays described above. Cell lines are shown in order of increasing sensitivity to cetuximab. As shown in FIG. 1, there is no correlation between EGFR level and sensitivity to cetuximab. Of the fourteen NSCLC cell lines tested, ChagoK1 and L2987 were the only two cell lines that consistently showed \geq 50% inhibition of cell proliferation at the IC50 concentration of cetuximab. Cell lines SW900, Calu6, SK-MES1, H838 and H661 showed significantly lower than 50% inhibition of cell proliferation at the doses of cetuximab that were tested. The remaining cell lines

LX1, H522, H441, H226, A549, SK-LU1 and H2347 showed no inhibition of cell proliferation at the doses of cetuximab that were tested. For the analysis, cell lines ChagoK1 and L2987 were defined as sensitive and the remaining twelve cell lines were defined as resistant.

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Gene Expression Profiling:

RNA for the NSCLC adenocarcinomas was purchased from a commercial vendor as described above. For the NSCLC cell lines, RNA was isolated from 50-70% confluent cells using the RNeasy kits (Qiagen, Valencia, CA). The quality of RNA was checked by measuring the 28S:18: ribosomal RNA ratio using an Agilent 2100 Bioanalyzer (Agilent Technologies, Rockville, MD). Concentration of total RNA was determined spectrophotometrically. 5 or 10 ug of total RNA was used to prepare biotinylated probes according to the Affymetrix Genechip Expression Analysis Technical Manual. Targets were hybridized to human HG-U133A gene chips according to the manufacturer's instructions. Data were preprocessed using the MAS 5.0 software (Affymetrix, Santa Clara, CA). The trimmed mean intensity for each chip was scaled to 1,500 to account for minor differences in global chip intensity so that the overall expression level for each sample is comparable.

20 Data Analysis

All 22,215 probes (gene sequences) present on the U133A chip were considered as potential predictive biomarkers. To restrict the analysis to gene sequences expressed in at least two of the twenty nine NSCLC tumors, gene sequences with Affymetrix MAS5.0 p> 0.04 in at least two tumors or cell lines were removed leaving 14,354 and 13,909 gene sequences, respectively (FIG. 2).

Next, to identify genes with variable expression in lung tumors (and therefore more likely to be able to correlate with variability in response to treatment), a variance metric (the Weighted spread (90-10) metric) (WSpread (90-10) metric) was used to calculate the variance of probe sets in the tumor and cell line expression profiling data.

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Weighted spread = $\underline{I \text{ 90th percentile - } I \text{ 10th percentile}}$

I = Signal intensity from expression profiling data

Gene sequences with a WSpread (90-10) metric < 30 were removed leaving 4167 gene sequences in the adenocarcinoma tumors (FIG. 3) and 4274 gene sequences in the cell lines (FIG. 4).

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Next, the same expression filter was applied to the remaining 4167 gene sequences using the NSCLC cell line data, resulting in 3572 gene sequences for analysis. This was followed by the application of the same variance metric filter leaving 2496 gene sequences for analysis. Of the 2496 gene sequences, 776 genes sequences ranked in the top 1000 in the cell line variance analysis. These 776 sequences were chosen for further statistical analysis. The 776 gene sequences were subjected to a two-sided unequal variance t-test using the resistance/sensitivity classifications of the cell lines described above (FIG. 1). 147 gene sequences showed a significantly different expression profile between the sensitive and resistant cell lines with a p-value of <0.05 (FIG. 5). Table 1 provides a list of the 147 gene sequences identified using the two-sided unequal variance T-test. These 147 gene sequences (probe sets) represent 124 biomarkers with regard to the Unigene Titles.

A variation of the gene filtering scheme illustrated in FIG. 1 was conducted and is illustrated in FIG. 2. In this scheme, 343 gene sequences ranked in the top 1000 in both the tumor and cell line variance analysis, a total of 343 out of the 776 genes sequences, were subjected to a two-sided unequal variance T-test. 59 gene sequences showed a significantly different expression profile between the sensitive and resistant cell lines with a p-value of <0.05. These 59 biomarkers are provided in Table 1 as the first 59 biomarkers, i.e., SEQ ID NOS:1-59 and 148-206.

EXAMPLE 2 – Experimental Validation of Biomarker Candidates: Cell line induction studies

Regulation by EGFR inhibitors in drug treated cell lines would lend additional support to the candidate biomarkers as being predictive of response. Induction

experiments were carried out in two sensitive cell lines ChagoK1 (sensitive to cetuximab and gefitinib) and L2987 (sensitive to cetuximab, resistant to gefitinib). Induction experiments were also carried out in four cell lines that were resistant to both EGFR inhibitors: A549 and H226 (EGFR+) and LX-1 and H522 (EGFR negative) cell lines.

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Cells were seeded in 6-well tissue culture dishes in DMEM supplemented with 10% FBS (Invitrogen, Carlsbad, CA). Twenty-four hours later the cells were switched to DMEM containing 0.5% FBS. The next day cells were treated with either 4µg/ml cetuximab or 1µM gefitinib. Twenty-four hours later cells were stimulated with 100ng/ml human recombinant epidermal growth factor EGF (Biosource International, Camarillo, CA) for 6 hours. The cells were lysed directly in the culture dish and RNA isolation was carried out using the RNeasy mini kit (Qiagen, Valencia, CA). Profiling was done on U133A GeneChips (Affymetrix, Santa Clara, CA). Data was analyzed using GeneChip® Expression Analysis software MAS 5.0 (Affymetrix, Santa Clara, California). Anova analysis of profiling data was done with PartekPro pattern recognition software (Partek, St. Charles, MS) using quantile normalized Affymetrix MAS 5.0 values for signal intensity.

Of the 147 probe sets examined, 21 probe sets representing 18 different biomarkers (provided below in Table 3) were highly regulated (Bonferroni p<0.05 in Anova analysis) upon EGFR inhibitor treatment and/or EGF stimulation in the sensitive cell lines.

TABLE 3 - Biomarkers Highly Regulated by EGFR Inhibitor Treatment and/or EGF Stimulation in the Sensitive Cell Lines

Unigene title and	Affymetrix Description	Affymetrix
SEQID NO:		Probe Set
DKK1: dickkopf	gb:NM_012242.1 /DEF=Homo sapiens	204602_at
homolog 1	dickkopf (Xenopus laevis) homolog 1 (DKK1),	
(LOC22943)	mRNA. /FEA=mRNA /GEN=DKK1	
	/PROD=dickkopf (Xenopus laevis) homolog 1	
SEQ ID NOS: 7	/DB_XREF=gi:7110718/UG=Hs.40499	
(DNA) and 154	dickkopf (Xenopus laevis) homolog 1	
(amino acid)	/FL=gb:AF127563.1 gb:AF177394.1	
	gb:NM 012242.1	

S100A9: S100	gb:NM_002965.2 /DEF=Homo sapiens S100	203535_at
calcium-binding	calcium-binding protein A9 (calgranulin B)	
protein A9	(S100A9), mRNA. /FEA=mRNA	
(LOC6280)	/GEN=S100A9 /PROD=S100 calcium-binding	
	protein A9 /DB_XREF=gi:9845520	
SEQ ID NOS: 10	/UG=Hs.112405 S100 calcium-binding protein	
(DNA) and 157	A9 (calgranulin B) /FL=gb:M26311.1	
(amino acid)	gb:NM_002965.2	
SFN: stratifin	Cluster Incl. X57348:H.sapiens mRNA (clone	33322_i_at
(LOC2810)	9112) /cds=(165,911) /gb=X57348 /gi=23939	
	/ug=Hs.184510 /len=1407	
SEQ ID NOS: 11		
(DNA) and 158		
(amino acid)		
PBEF: pre-B-cell	Consensus includes gb:BF575514 /FEA=EST	217738_at
colony-enhancing	/DB XREF=gi:11649318	_
factor isoform a	/DB_XREF=est:602133090F1	
(LOC10135)	/CLONE=IMAGE:4288079 /UG=Hs.239138	
	pre-B-cell colony-enhancing factor	
SEQ ID NOS: 36	/FL=gb:U02020.1 gb:NM 005746.1	
(DNA) and 183		
(amino acid)		
SERPINE2:	Consensus includes gb:AL541302 /FEA=EST	212190_at
plasminogen activator	/DB XREF=gi:12872241	_
inhibitor type 1,	/DB XREF=est:AL541302	
member 2	/CLONE=CS0DE006YI10 (5 prime)	
(LOC5270)	/UG=Hs.21858 trinucleotide repeat containing	
	3	
SEQ ID NOS: 38		
(DNA) and 185		
(amino acid)		
SFN: stratifin	Cluster Incl. X57348:H.sapiens mRNA (clone	33323_r_at
(LOC2810)	9112) /cds=(165,911) /gb=X57348 /gi=23939	
	/ug=Hs.184510 /len=1407	
SEQ ID NOS: 41		
(DNA) and 188		
(amino acid)		
IL8: interleukin 8	gb:AF043337.1 /DEF=Homo sapiens	211506_s_at
(LOC3576)	interleukin 8 C-terminal variant (IL8) mRNA,	
,	complete cds. /FEA=mRNA /GEN=IL8	
SEQ ID NOS: 44	/PROD=interleukin 8 C-terminal variant	
(DNA) and 191	/DB_XREF=gi:12641914 /UG=Hs.624	
(amino acid)	interleukin 8 /FL=gb:AF043337.1	

CTSC: cathepsin C	gb:NM 001814.1 /DEF=Homo sapiens	201487 at
isoform a	cathepsin C (CTSC), mRNA. /FEA=mRNA	_
preproprotein	/GEN=CTSC /PROD=cathepsin C	
(LOC1075)	/DB XREF=gi:4503140 /UG=Hs.10029	
(2001073)	cathepsin C /FL=gb:NM_001814.1	
SEQ ID NOS: 46	Surropoint Cit E gon tin_coror in	
(DNA) and 193		
(amino acid)		
TXNIP: thioredoxin	Consensus includes gb:AA812232 /FEA=EST	201008 s at
	/DB XREF=gi:2881843	201006_s_at
interacting protein	/DB_XREF=g1.2881845 /DB_XREF=est:ob84h09.s1	
(LOC10628)	<u> </u>	,
	/CLONE=IMAGE:1338113 /UG=Hs.179526	
SEQ ID NOS: 50	upregulated by 1,25-dihydroxyvitamin D-3	
(DNA) and 197	/FL=gb:NM_006472.1 gb:S73591.1	
(amino acid)		-10-500
SAT:	gb:M55580.1 /DEF=Human	210592_s_at
spermidine/spermine	spermidinespermine N1-acetyltransferase	
N1-acetyltransferase	mRNA, complete cds. /FEA=mRNA	
(LOC6303)	/GEN=spermidinespermine N1-	
	acetyltransferase /PROD=spermidinespermine	
SEQ ID NOS: 54	N1-acetyltransferase /DB_XREF=gi:338335	
(DNA) and 201	/UG=Hs.28491 spermidinespermine N1-	,
(amino acid)	acetyltransferase /FL=gb:M55580.1	
TXNIP: thioredoxin	gb:NM 006472.1 /DEF=Homo sapiens	201010 s at
interacting protein	upregulated by 1,25-dihydroxyvitamin D-3	
(LOC10628)	(VDUP1), mRNA. /FEA=mRNA	
(20010020)	/GEN=VDUP1 /PROD=upregulated by 1,25-	
SEQ ID NOS: 57	dihydroxyvitamin D-3 /DB_XREF=gi:5454161	
(DNA) and 204	/UG=Hs.179526 upregulated by 1,25-	
(amino acid)	dihydroxyvitamin D-3 /FL=gb:NM_006472.1	
(diffine dela)	gb:S73591.1	
TENS1: tensin-like	gb:NM 022748.1 /DEF=Homo sapiens	217853_at
SH2 domain-	hypothetical protein FLJ13732 similar to tensin	217035_40
containing 1	(FLJ13732), mRNA. /FEA=mRNA	
(LOC64759)	/GEN=FLJ13732 /PROD=hypothetical protein	,
(LOC04739)	FLJ13732 similar to tensin	
SEO ID NOS. 66	/DB XREF=gi:12232408/UG=Hs.12210	
SEQ ID NOS: 66	hypothetical protein FLJ13732 similar to tensin	
(DNA) and 213	/FL=gb:NM_022748.1	
(amino acid)		202602 a at
STK17A:	Consensus includes gb:AW194730 /FEA=EST	202693_s_at
serine/threonine	/DB_XREF=gi:6473630	
kinase 17a	/DB_XREF=est:xn43d11.x1	
(apoptosis-inducing)	/CLONE=IMAGE:2696469 /UG=Hs.9075	
(LOC9263)	serinethreonine kinase 17a (apopto sis-inducing)	
	/FL=gb:AB011420.1 gb:NM_004760.1	
SEQ ID NOS: 69	t .	
(DNA) and 216		
(amino acid)		

TUBB-5: tubulin	gb:BC002654.1 /DEF=Homo sapiens, Similar	209191 at
beta-5 (LOC84617)	to tubulin, beta, 4, clone MGC:4083, mRNA,	200101_46
Deta 5 (Ecco (017)	complete cds. /FEA=mRNA /PROD=Similar to	
SEQ ID NOS: 84	tubulin, beta, 4 /DB XREF=gi:12803638	
(DNA) and 231	/UG=Hs.274398 Homo sapiens, Similar to	
(amino acid)	tubulin, beta, 4, clone MGC:4083, mRNA,	
(diffile deld)	complete cds /FL=gb:BC002654.1	
TYMS: thymidylate	gb:NM 001071.1 /DEF=Homo sapiers	202589 at
synthetase	thymidylate synthetase (TYMS), mRNA.	202367_at
(LOC 7298)	/FEA=mRNA /GEN=TYMS	
(LOC 7298)	/PROD=thymidylate synthetase	
SEQ ID NOS: 85	/DB XREF=gi:4507750 /UG=Hs.82962	
(DNA) and 232	thymidylate synthetase /FL=gb:BC002567.1	
(amino acid)	gb:NM 001071.1	
RAI14: retinoic acid	gb:NM 01577.1 /DEF=Homo sapiens novel	202052 s at
induced 14	retinal pigment epithelial gene (NORPEG),	202032_8_at
1	mRNA. /FEA=mRNA /GEN=NORPEG	
(LOC26064)		
SEO ED NOS: 07	/PROD=DKFZP564G013 protein	
SEQ ID NOS: 97	/DB_XREF=gi:13470085 /UG=Hs.15165 novel	
(DNA) and 244	retinal pigment epithelial gene	
(amino acid)	/FL=gb:NM_015577.1 gb:AF155135.1	010077 -4
CALD1: caldesmon 1	Consensus includes gb:AL583520 /FEA=EST	212077_at
isoform 3 (LOC800)	/DB_XREF=gi:12952562	
GEO ED NIOG 106	/DB_XREF=est:AL583520	
SEQ ID NOS: 106	/CLONE=CS0DC024YE13 (5 prime)	
(DNA) and 253	/UG=Hs.182183 Homo sapiens mRNA for	
(amino acid)	caldesmon, 3 UTR	202760
PALM2: paralemmin	gb:NM_007203.1 /DEF=Homo sapiens A	202760_s_at
2 (LOC114299)	kinase (PRKA) anchor protein 2 (AKAP2),	
	mRNA. /FEA=mRNA /GEN=AKAP2	
SEQ ID NOS: 115	/PROD=A kinase (PRKA) anchor protein 2	,
(DNA) and 262	/DB_XREF=gi:6005708 /UG=Hs.423 22 A	
(amino acid)	kinase (PRKA) anchor protein 2	
	/FL=gb:AB023137.1 gb:NM_007203.1	-1000
TPM1: tropomyosin 1	gb:Z24727.1 /DEF=H.sapiens tropomyosin	210986_s_at
(alpha) (LOC7168)	isoform mRNA, complete CDS. /FEA=mRNA	
	/PROD=tropomyosin isoform	
SEQ ID NOS: 125	/DB_XREF=gi:854188 /UG=Hs.77899	
(DNA) and 272	tropomyosin 1 (alpha) /FL=gb:Z24727.1	
(amino acid)		
TPM1: tropomyosin 1	gb:M19267.1 /DEF=Human tropomyosin	210987_x_at
(alpha) (LOC7168)	mRNA, complete cds. /FEA=mRNA	
	/DB_XREF=gi:339943 /UG=Hs.77899	
SEQ ID NOS: 137	tropomyosin 1 (alpha) /FL≈gb:M19267.1	
(DNA) and 284		
(amino acid)		

TUBB: tubulin, beta	gb:NM_001069.1 /DEF=Homo sapiens tubulin,	204141_at
polypeptide	beta polypeptide (TUBB), mRNA.	
(LOC7280)	/FEA=mRNA /GEN=TUBB /PROD=tubulin,	
	beta polypeptide /DB XREF=gi:4507728	
SEQ ID NOS: 147	/UG=Hs.179661 tubulin, beta polypeptide	
(DNA) and 294	/FL=gb:BC001194.1 gb:NM 001069.1	
(amino acid)		

It appears that these biomarkers are likely to be directly or indirectly involved in the EGFR signaling pathway, based on their expression modulation by EGF and / or EGFR inhibitor treatment.

EXAMPLE 3 - Experimental Validation of Biomarker Candidates: Drug treatment

Regulation by EGFR inhibitors in lung xenograft models would lend additional support to the candidate markers, as being predictive of response. Drug treatment experiments were carried out in the L2987 (sensitive to cetuximab and gefitinib), A549 (borderline sensitive to cetuximab and gefitinib), and LX1 (resistant to cetuximab and gefitinib) lung xenograft models.

In Vivo Antitumor Testing

studies in lung xenograft models

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Tumors were propagated in nude mice as subcutaneous (sc) transplants using tumor fragments obtained from donor mice. Tumor passage occurred approximately every two to four weeks. Tumors were then allowed to grow to the pre-determined size window (usually between 100-200 mg, tumors outside the range were excluded) and animals were evenly distributed to various treatment and control groups. Animals were treated with cetuximab (1 mg/mouse, q3d X 10, 14; ip) or gefitinib (200mg/kg, q1dX14, 14; po). Treated animals were checked daily for treatment related toxicity/mortality. Each group of animals was weighed before the initiation of treatment (Wt1) and then again following the last treatment dose (Wt2). The difference in body weight (Wt2-Wt1) provided a measure of treatment-related toxicity. Tumor response was determined by measurement of tumors with a caliper twice a week, until the tumors reached a predetermined target size of 1 gm or became necrotic. Tumor weights (mg) were estimated from the formula:

Tumor weight = $(length \times width^2)/2$

Antitumor activity was determined in terms of primary tumor growth inhibition. This was determined in two ways: (i) calculating the relative median tumor weight (MTW) of treated (T) and control (C) mice at various time points (effects were expressed as %T/C); and (ii) calculating the tumor growth delay (T-C value), defined as the difference in time (days) required for the treated tumors (T) to reach a predetermined target size compared to those of the control group (C). Statistical evaluations of data were performed using Gehan's generalized Wilcoxon test for comparisons of time to reach tumor target size (Gehan 1965). Statistical significance was declared at p < 0.05. Antitumor activity was defined as a continuous MTW $\%T/C \le 50\%$ for at least 1 tumor volume doubling time (TVDT) any time after the start of treatment, where TVDT (tumor volume doubling time) = median time (days) for control tumors to reach target size – median time (days) for control tumors to reach half the target size. In addition, treatment groups had to be accompanied by a statistically significant tumor growth delay (T-C value) (p < 0.05) to be termed active.

Treated animals were checked daily for treatment related toxicity/mortality. When death occurred, the day of death was recorded. Treated mice dying prior to having their tumors reach target size were considered to have died from drug toxicity. No control mice died bearing tumors less than target size. Treatment groups with more than one death caused by drug toxicity were considered to have had excessively toxic treatments and their data were not included in the evaluation of the compound's antitumor efficacy.

Drug treatment experiments

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L2987 and A549 xenograft animals were dosed with a single dose of either (1) 1 mg/mouse cetuximab, ip; (2) 250mg/kg gefitinib, po; (3) PEG400/H₂O vehicle, po or 4) PBS vehicle, ip. Each dose was given to three independent mice. At 3h and 24h post-treatment the animals were sacrificed and tumors were excised and immediately placed into RNA*later* solution (Oiagen, Valencia, CA).

RNA was isolated from the tumors using the RNeasy kits (Qiagen, Valencia, CA). The quality and concentration of total RNA was determined as described previously. Profiling was done on U133A GeneChips (Affymetrix, Santa Clara, CA).

Data was analyzed using GeneChip® Expression Analysis software MAS 5.0 (Affymetrix, Santa Clara, California). Anova analysis of profiling data was done with PartekPro pattern recognition software (Partek, St. Charles, MS) using quantile normalized Affymetrix MAS5.0 values for signal intensity.

Out of 147 probesets examined, 4 probesets representing 3 genes are significantly regulated (p<0.005 in Anova analysis) upon EGFR inhibitor treatment in the sensitive L2987 xenograft but not in the borderline sensitive A549 xenograft. The three genes are jumping translocation breakpoint (JTB), 3-phosphoadenosine 5-phosphosulfate synthase 2 (PAPSS2) and serine protease inhibitor, Kunitz type 1 (SPINT1). It appears that these biomarkers are likely to be directly or indirectly involved in the EGFR signaling pathway, based on their expression modulation by EGFR inhibitor treatment.

EXAMPLE 4 - Immunohistochemistry (IHC) assays in clinical samples

Of the 147 probe sets identified preclinically, S100A9 (Calgranulin B) was
chosen to examine whether there was any correlation between expression of a
particular protein in the clinical samples and Best Clinical Response data.

Basic IHC Method

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Formalin-fixed, paraffin-embedded tissues were available on slides in 5 µm sections. The sections were deparaffinized with standard xylene and hydrated through graded alcohols into water. Antigen retrieval was performed using proteinase K. Staining was done at room temperature on an automatic staining workstation TechMate 1000 (BioTek Solutions/Ventana Medical Systems, Tucson, AZ) by using the Envision peroxidase mouse system (DakoCytomation, Carpinteria, CA). Slides were placed three times for 2.5 minutes each in a hydrogen peroxide blocking medium and then allowed to react with mouse anti-human Calgranulin B monoclonal antibody (Bachem Biomedical, Germany) for 60 minutes. Immunodetection was performed with the Envision system by placing slides three times for 5 minutes each in diaminobenzidine (DAB) chromogen substrate. Counterstaining with hematoxylin for 1 minute was the final step. After staining, slides were dehydrated through an alcohol series to absolute ethanol followed by xylene rinses. Slides were permanently

coverslipped with glass coverslips and permount medium. Slides were examined under a microscope to assess staining. Positive staining is indicated by the presence of a dark brown chromogen (DAB-Horse Radish Peroxidase reaction product). Hematoxylin counterstain provides a blue nuclear stain to assess cell and tissue morphology. Appropriate positive and negative controls were used. The slides were viewed randomly, without clinical data, by two independent evaluators and scored. A simple scoring system was used to reflect whether a tissue is positive or negative for the marker and to indicate the relative level of staining. A scoring scheme of negative, low, moderate or high was used to indicate the relative percentage of tumor cells staining within the tissues (FIG. 7). The scoring system simply provides an indication of relative expression of a target from tissue to tissue.

Clinical materials and criteria for response

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Formalin-fixed paraffin embedded lung tumor slides were obtained from patients enrolled in a phase II trial of cetuximab. In this trial, cetuximab was used as a single agent therapy for recurrent non-small-cell lung cancer patients (unpublished). The best overall response was recorded from the start of the treatment until disease progression or recurrence. Assessment of response was performed using the RECIST criteria (Response Evaluation Criteria in Solid Tumors, Tsuchida and Therasse, 2001). A partial response (PR) described at least a 30% decrease in the sum of the longest diameter (LD) of target lesions, taking as reference the baseline sum LD. Progressive disease (PD) referred to a 20% or greater increase in the sum of the LD of target lesions, taking as reference the smallest sum LD recorded since the treatment started or the appearance of new lesions. Stable Disease (SD) was used to describe neither sufficient shrinkage to qualify for PR nor sufficient increase to qualify for PD.

Calgranulin B IHC assay on clinical FFPET slides

Calgranulin B IHC assay was performed on FFPET slides from 39 patients enrolled in the phase II trial of cetuximab in recurrent NSCLC patients (Table 4). Of the 39 patients, 10 were excluded from further analysis because there was no detectable tumor specimen on the slide. The remaining 29 patients that were scored for Calgranulin B staining comprised of 2 PR, 12 SD and 15 PD non-responders

based on the clinical response data. The 39 samples used in this IHC analysis were derived from patients for whom tissue samples were available and from whom an informed consent could be obtained. It should be noted that the response data shown here may not reflect the response rate in the entire study.

Of the 29 patients' slides, 22 were scored as 0, 3 were scored as 0.5+, 3 were scored as 1+ and 1 slide was scored as 2+. Overall 24 % of the patients tested were positive for Calgranulin B staining (Table 4).

TABLE 4 - IHC Assay Results

PROGRESSIVE DISEASE		DISEASE STABILIZATION			
	Best			Best	
	Clinical			Clinical	
ID	Response	\mathbf{IHC}	ID	Response	IHC
L8	PD	negative	L10	SD	negative
L11	PD	negative	L13	SD	positive
L12	PD	positive	L40	SD	negative
L14	PD	negative	L24	SD	negative
L15	PD	negative	L27	SD	positive
L18	PD	negative	L47	SD	positive
L20	PD	negative	L28	SD	negative
L41	PD	negative	L3	SD	negative
L42	PD	negative	L4	SD	negative
L44	PD	negative	L6	SD	positive
L16	PD	negative	L34	SD	negative
L5	PD	negative	L39	SD	positive
L33	PD	negative	L1	PR	positive
L37	PD	negative	L2	PR	negative
L23B	PD	negative			

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The results are summarized in Table 5 below.

TABLE 5 - IHC Assay Results Summary

	_	# non- responders
Calgranulin B +	6	1
Calgranulin B -	9	13

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Of the 7 patients that were Calgranulin B positive, 6 had disease stabilization and 1 was a non-responder having progressive disease (Table 5). The sensitivity of the

assay to identify potential responders is 40% [6/ (6+9)] and the specificity is 93% [13/ (13+1)].

The positive predictive value of a Calgranulin B IHC assay to identify potential responders is 86% [6/(6+1)] and the negative predictive value = 59% [13/(13+9)], {Chi square p value = 0.03}.

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Although the data set is small, these results indicate a trend for Calgranulin B positive patients to have disease stabilization.

EXAMPLE 5 - PRODUCTION OF ANTIBODIES AGAINST THE BIOMARKERS

Antibodies against the biomarkers can be prepared by a variety of methods. For example, cells expressing a biomarker polypeptide can be administered to an animal to induce the production of sera containing polyclonal antibodies directed to the expressed polypeptides. In one aspect, the biomarker protein is prepared and isolated or otherwise purified to render it substantially free of natural contaminants, using techniques commonly practiced in the art. Such a preparation is then introduced into an animal in order to produce polyclonal antisera of greater specific activity for the expressed and isolated polypeptide.

In one aspect, the antibodies of the invention are monoclonal antibodies (or protein binding fragments thereof). Cells expressing the biomarker polypeptide can be cultured in any suitable tissue culture medium, however, it is preferable to culture cells in Earle's modified Eagle's medium supplemented to contain 10% fetal bovine serum (inactivated at about 56 °C), and supplemented to contain about 10 g/l nonessential amino acids, about 1,00 U/ml penicillin, and about 100 μ g/ml streptomycin.

The splenocytes of immunized (and boosted) mice can be extracted and fused with a suitable myeloma cell line. Any suitable myeloma cell line can be employed in accordance with the invention, however, it is preferable to employ the parent myeloma cell line (SP2/0), available from the ATCC (Manassas, VA). After fusion, the resulting hybridoma cells are selectively maintained in HAT medium, and then cloned by limiting dilution as described by Wands et al. (1981, *Gastroenterology*, 80:225-232). The hybridoma cells obtained through such a selection are then assayed

to identify those cell clones that secrete antibodies capable of binding to the polypeptide immunogen, or a portion thereof.

Alternatively, additional antibodies capable of binding to the biomarker polypeptide can be produced in a two-step procedure using anti-idiotypic antibodies. Such a method makes use of the fact that antibodies are themselves antigens and, therefore, it is possible to obtain an antibody that binds to a second antibody. In accordance with this method, protein specific antibodies can be used to immunize an animal, preferably a mouse. The splenocytes of such an immunized animal are then used to produce hybridoma cells, and the hybridoma cells are screened to identify clones that produce an antibody whose ability to bind to the protein-specific antibody can be blocked by the polypeptide. Such antibodies comprise anti-idiotypic antibodies to the protein-specific antibody and can be used to immunize an animal to induce the formation of further protein-specific antibodies.

15 EXAMPLE 6 - IMMUNOFLUORESCENCE ASSAYS

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The following immunofluorescence protocol may be used, for example, to verify EGFR biomarker protein expression on cells or, for example, to check for the presence of one or more antibodies that bind EGFR biomarkers expressed on the surface of cells. Briefly, Lab-Tek II chamber slides are coated overnight at 4 °C with 10 micrograms/milliliter (µg/ml) of bovine collagen Type II in DPBS containing calcium and magnesium (DPBS++). The slides are then washed twice with cold DPBS++ and seeded with 8000 CHO-CCR5 or CHO pC4 transfected cells in a total volume of 125 µl and incubated at 37 °C in the presence of 95% oxygen / 5% carbon dioxide.

The culture medium is gently removed by aspiration and the adherent cells are washed twice with DPBS++ at ambient temperature. The slides are blocked with DPBS++ containing 0.2% BSA (blocker) at 0-4 °C for one hour. The blocking solution is gently removed by aspiration, and 125 µl of antibody containing solution (an antibody containing solution may be, for example, a hybridoma culture supernatant which is usually used undiluted, or serum/plasma which is usually diluted, e.g., a dilution of about 1/100 dilution). The slides are incubated for 1 hour at 0-4 °C. Antibody solutions are then gently removed by aspiration and the cells are

washed five times with 400 μ l of ice cold blocking solution. Next, 125 μ l of 1 μ g/ml rhodamine labeled secondary antibody (e.g., anti-human IgG) in blocker solution is added to the cells. Again, cells are incubated for 1 hour at 0-4 °C.

The secondary antibody solution is then gently removed by aspiration and the cells are washed three times with 400 µl of ice cold blocking solution, and five times with cold DPBS++. The cells are then fixed with 125 µl of 3.7% formaldehyde in DPBS++ for 15 minutes at ambient temperature. Thereafter, the cells are washed five times with 400 µl of DPBS++ at ambient temperature. Finally, the cells are mounted in 50% aqueous glycerol and viewed in a fluorescence microscope using rhodamine filters.

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CLAIMS:

What is claimed is:

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1. A method for identifying a mammal that will respond therapeutically to a method of treating cancer comprising administering an EGFR modulator, wherein the method comprises:

- (a) measuring in the mammal the level of at least one biomarker selected from the biomarkers of Table 1;
 - (b) exposing a biological sample from said mammal to the EGFR modulator;
- (c) following the exposing of step (b), measuring in said biological sample the level of the at least one biomarker,

wherein a difference in the level of the at least one biomarker measured in step (c) compared to the level of the at least one biomarker measured in step (a) indicates that the mammal will respond therapeutically to said method of treating cancer.

- 2. A method for identifying a mammal that will respond therapeutically to a method of treating cancer comprising administering an EGFR modulator, wherein the method comprises:
 - (a) exposing a biological sample from the mammal to the EGFR modulator;
- (b) following the exposing of step (a), measuring in said biological sample the level of the at least one biomarker selected from the biomarkers of Table 1,
- wherein a difference in the level of the at least one biomarker measured in step (b), compared to the level of the at least one biomarker in a mammal that has not been exposed to said EGFR modulator, indicates that the mammal will respond therapeutically to said method of treating cancer.

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Identification Scheme of Table 1 Biomarkers

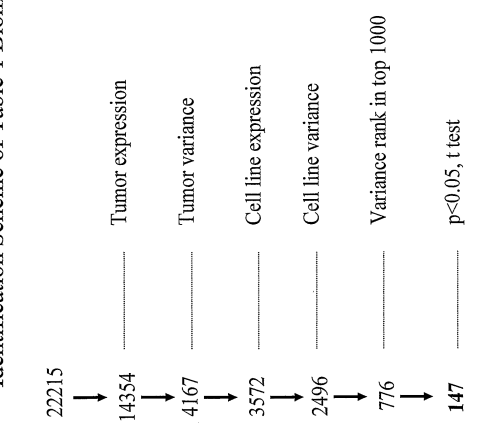


FIG. 1

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Identification Scheme of Table 2 Biomarkers

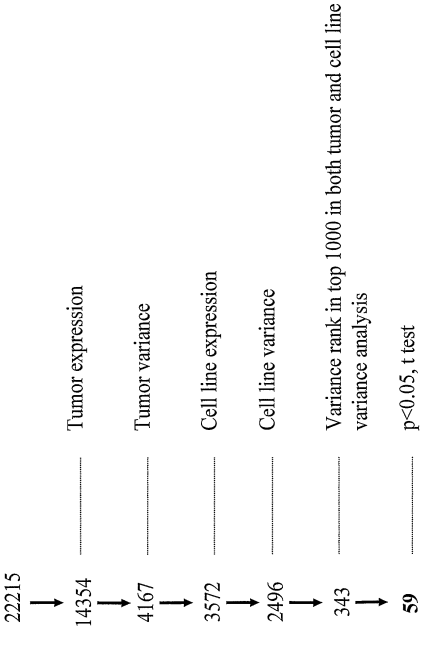


FIG. 2

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EGFR Expression in NSCLC Cell Lines

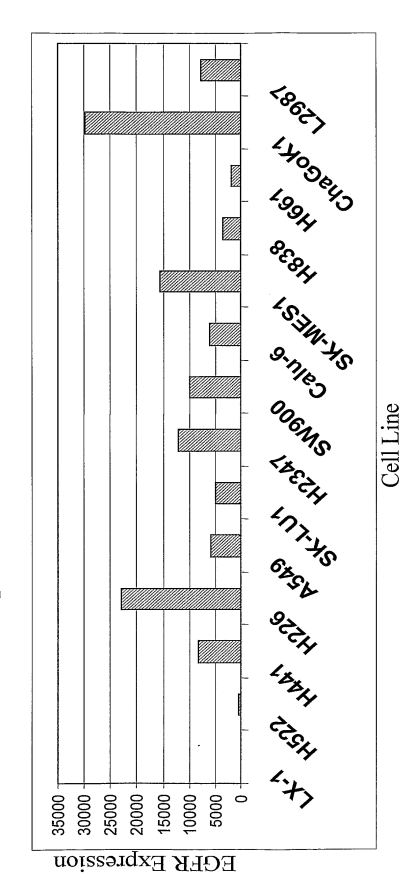


FIG. 3

Variance Analysis of Expression Profiles

29 Adenocarcinoma Tumors

22,215 probe set Affymetrix GeneChip® HG-U133A array profiles of 29 lung adenocarcinoma tumors

- 14,354 probe sets present (p \leq 0.04) in at least 2 samples
- Calculate variance and rank probe sets

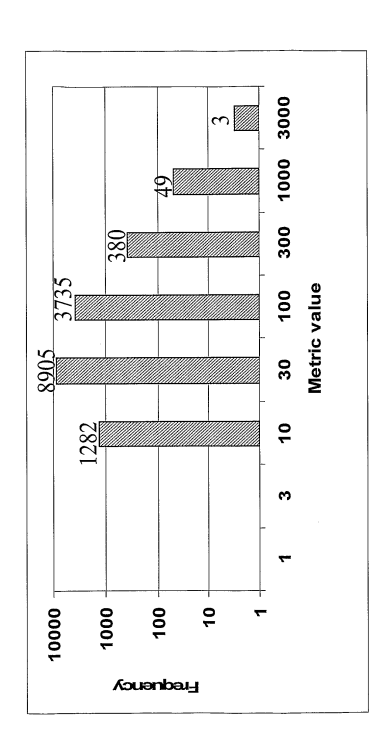
14 NSCLC Cell Lines

- GeneChip® HG-U133A array profiles of 14 NSCLC cell lines
- 13,909 probe sets present ($p \le 0.04$) in at least 2 cell lines
- Calculate variance and rank probe sets

FIG. 4

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Variance Metric Distribution of Probe sets

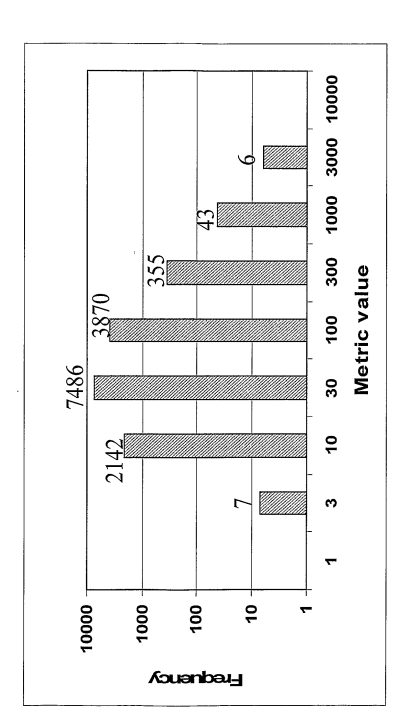


Adenocarcinoma Tumors - 4167 probe sets of WSpread(90-10) ≥ 30

FIG. 5

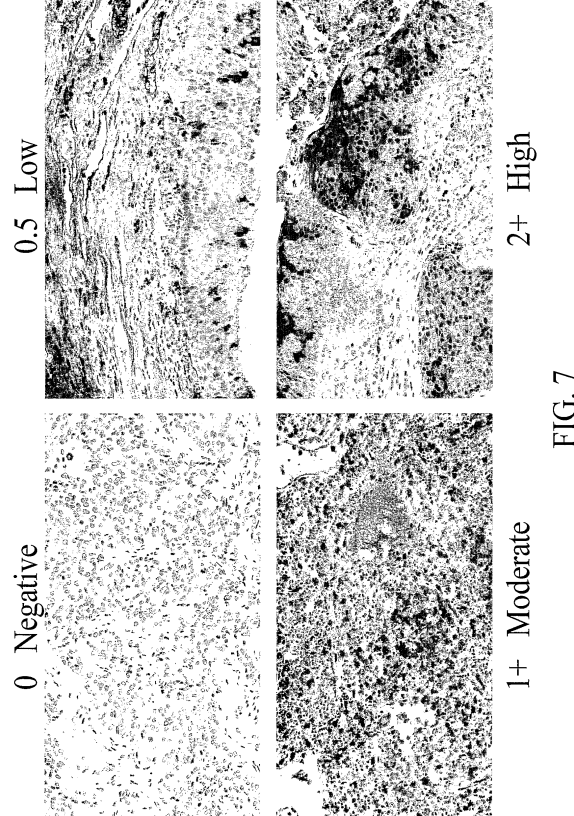
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Variance Metric Distribution of Probe sets



Cell Lines - 4274 probe sets of WSpread(90-10) ≥ 30

FIG. 6



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